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Impact of Dual Calcium and Manganese Substitution of La-deficient perovskites on structural and related Catalytic Properties : Future opportunities in next Three-Way-Catalyst Generation?

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Abstract

The impact of La-deficiency and substitution by calcium and manganese in LaFeO₃ has been investigated in Three-Way-Catalysis. Despite significant structural distortions, predominantly induced by calcium substitution, no significant phase segregation occurs. On the other hand, bulk and surface characterization revealed the stabilization of Mn⁴⁺ and Fe⁴⁺ in different extent. Kinetic analysis of CO and propene oxidation reactions and the NO reduction reaction showed that La_{0.7}Fe_{0.8}Mn_{0.2}O₃ exhibits the best performance. By contrast, a detrimental effect of calcium more pronounced for propene oxidation is observed. It was found that catalytic properties are neither related to bulk properties nor to the extent of specific surface area but are mostly governed by the redox couple Mn⁴⁺/Mn³⁺. The formation of Fe⁴⁺ improving with partial substitution of La by Ca in A-site was not found as a determining criterion.

Keywords: Perovskite, Three-Way-Catalyst, La-deficient La_{1-x}FeO₃, Dual calcium manganese substitution, CO and propene oxidation, NO reduction.

1. Introduction

Perovskites (ABO₃) structures have been utilized in many oxidative catalytic applications and catalytic end-of-pipe technologies to treat atmospheric pollutants from Diesel exhaust, i.e. Diesel oxidation Catalysts, NO_x Storage Reduction catalysts [1-4]. Among the different compositions previously studied, stoichiometric LaMnO₃ [1,3] and none-stoichiometry La_xMnO₃ structures [5] attracted the interest of several investigators. As example, Chen et al. [5] reported improved performance for NO oxidation to NO₂ on La_{0.9}MnO₃ ascribed to higher formation of Mn⁴⁺. These authors explained this gain in catalytic properties to the formation of more active oxygen species bonded to Mn⁴⁺. The catalytic properties of LaMn_{0.9}O₃ were also examined for methane combustion with improved performances related to specific surface area enhancement jointly to higher density of defective sites [6]. Manganites have been extensively studied in oxidation reactions for light alkanes (C₂-C₇) and alkenes. [7,8].

The stabilization of Mn^{4+} instead of Mn^{3+} occurs even for stoichiometric LaMnO₃ composition. In that case, the excess of positive charges will be counterbalanced by an oxygen overstoichiometry. As reported elsewhere [9], perovskite structures can accommodate a wide variety of cations in B-site with mixed valency in case of non-stoichiometry of B or A cations or partial substitution with cations having different oxidation states. By way of illustration, Onrubia et al. [1] found that La³⁺ substitution by Sr²⁺ in Sr-doped LaCoO₃ induces the creation of oxygen vacancies with unchanged oxidation state for cobalt. On the other hand, these authors observed that the same substitution in Sr-doped LaMnO₃ preferentially leads to an increase of the oxidation state of manganese shifting to Mn⁴⁺. For dual site substitutions, such as La³⁺ by Ca²⁺ in A-site and Co³⁺ by Mg²⁺ in B-site of parent LaCoO₃ material, a shift of the oxidation state of Co from +III to +IV is observed leading to a rate enhancement in toluene oxidation. Such gain in catalytic activity has been explained from a preferential suprafacial mechanism. rather than intrafacial mechanism [9].

Starting from stoichiometric LaFeO₃ and La-deficient La_{0.7}FeO₃ as benchmark compositions, single and dual substitution have been investigated in this study. Particular attention was paid to change in structural and electronic features induced by such substitutions which can lead to the stabilization of manganese or iron in unusual +IV oxidation state [10]. Accordingly, two redox couples, Fe^{4+}/Fe^{3+} and Mn^{4+}/Mn^{3+} can coexist and could influence the resulting catalytic properties. Leontiou et al. [11] found a greater stabilization of Fe^{4+} in Sr-doped LaFeO₃ from Mössbauer spectroscopy. These authors observed that the highest activity in NO reduction by CO was obtained on the perovskites having the lowest amount of Fe^{4+} and the highest amount of Fe^{3+} . Zhu and Thomas [12] reported that understanding the impact of B-site substitution on the overall performance is not trivial due to the coexistence of two redox systems which can be equally involved in NO decomposition. The identification of the most efficient redox couple remains empirical and is not formally related to a permanent oxidation state or the coexistence of different oxidation states [12].

Despite higher scarce examples, the potential interest of perovskites has been already tackled for three-way-catalysts but in simulated conditions which are often far from the typical compositions of the automotive exhaust gas. Previous investigations showed that LaMnO₃ catalyst can be a good candidate for CO [13] and hydrocarbon oxidation [14] as well as NO reduction [16-18]. Improved performance in NO reduction by propene have been reported on Cu-substituted LaMnO₃ perovskite in stoichiometric or slightly oxidizing conditions related to adsorbed ad-NO_x intermediates through the oxidation of NO by adsorbed oxygen species on anionic vacancies. On the other hand, a detrimental effect is observed with a rise in gaseous oxygen concentration because an excessive surface concentration of active oxygen species favors the preferential oxidation of propene [15-17].

Iron perovskites were selected as parent materials because of their improved thermal stability giving rise to better durability [18]. The impact of lanthanum substitution by calcium and iron

by manganese on the catalytic performance in three-way conditions has been investigated taking the occurrence of possible phase segregation into account that can improve/deteriorate activity and stability as well as modify the balance between reductive and oxidative properties [10].

2. Experimental

2.1. Catalyst preparation

Sol-gel method earlier described [19] was implemented for the preparation of parent and dual substituted perovskites according to a citrate route. Lanthanum, iron, manganese nitrate precursor salts were dissolved with citric acid (AC) in deionized water with CA/A+B =1. The solution thus obtained was subjected to evaporation in a rotary evaporator at 60°C with a rotating velocity of 30 rpm up to the formation of a gel. After drying in a static oven at 80°C for 48 h, the dried sample was calcined in air at 600°C for 8 h with a heating temperature dT/dt = 2°C/min.

2.2. Physicochemical characterization

X-ray diffraction (XRD) measurements were performed on a Bruker AXS D8 diffractometer (Cu K α source, $\lambda = 0.154$ nm). XRD patterns were recorded in the 2θ range of 10-80 degree. Lattice parameters and unit cell volumes were adjusted from the Rietveld refinement method. Crystallite size was estimated by the Williamson-Hall approach. Standard deviations were calculated by Bérar correction.

Ex situ Mössbauer spectra were recorded at room temperature using a ⁵⁷Co (Rh) source. Lorentzian shape peaks were assumed for data fitting. Pure iron foil was used as reference for the calculation of hyperfine field (HF), isomer shift (IS) and quadrupole splitting (QS). Scanning Electron Microscopy (SEM) micrographs were recorded on a Hitachi S-4700 Cold Field Emission Gun Scanning Electron Microscope operating at an acceleration voltage of 15 kV. Energy-dispersive X-ray spectroscopy (EDS) was used for elemental analysis. Oxford EDS system was employed for these analyzes. The volume probed by this technique was 1 μ m³.

H₂-Temperature-Programmed Reduction (H₂-TPR) experiments were carried out on a Micromeritics AutoChem 2920 analyzer. 50 mg samples, subjected to 50 mL/min flowrate of 5 vol.% H₂ in Ar, were heated to 1000°C with a heating rate of 5 °C/min. Hydrogen consumption was controlled by a thermal conductivity detector. O₂-Temperature-Programmed-Desorption (O₂-TPD) experiments were performed on the same apparatus with the same mass of catalyst. The sample was heated under He up to 1000°C with dT/dt = 10 °C/min. Oxygen concentration in the gas phase released from the catalyst surface during the ramp of temperature was measured by gas chromatography and mass spectrometery (m/z = 32). Prior to O₂-TPD experiments, the samples were systematically pre-heated in pure oxygen at 600°C for 1 h.

Porosity measurements on a Micromeritics Tristar analyzer were carried out -196°C. Prior to nitrogen physisorption, the samples were outgassed under vacuum for 4 h. Specific surface area (SSA) was determined according to Brunauer-Emmett-Teller (BET) theory, while the Barret-Joyner-Halenda (BJH) theory led to pore size distribution.

X-ray photoelectron spectra were recorded on a Kratos Axis Ultra spectrometer equiped with a monochromatized Al K α source. Binding Energy (B.E.) values were corrected compared to

the C1s core level set at 284.8 eV. Casa XPS software was used for spectral decomposition accounting for mixed Gaussian/Lorentzian (70/30) line profile.

2.3. Catalytic measurements

Temperature-Programmed-Reaction experiments (TPR) were performed at atmospheric pressure according to the same protocol described elsewhere [10]. A fixed bed flow reactor was fed with 200 mg catalysts, with average grain size of 150 μ m, mixed with the same mass of SiC. The total inlet volumetric flowrate (Q₀) at 12 L h⁻¹, corresponded to Q₀/W of 60,000 mL h⁻¹ g⁻¹. The protocol depicted in Fig. S1 in Supplementary Information (SI) included 4 successive TPR experiments performed in stoichiometric, lean and rich conditions with gas mixture compositions in Table S1 in SI. The outlet NO, NO₂, N₂, N₂O, O₂, H₂, CO, C₃H₆ and C₃H₈ concentrations were controlled by a Thermo Scientific Model 42i-HL chemiluminescence NO_x analyzer and an Agilent 490 Micro Gas Chromatography. Rate calculations were detailed in SI assuming the mass balance equation of a plug flow reactor in agreement with Eqs. (S1)-(S3) in SI. Preliminary verifications showed that mass transfer limitations should not occur significantly for conversion lower than 0.4 in agreement with the boundary conditions given by Eq. (S4) in SI.

3. Results and discussion

3.1. Induced effect of single and dual calcium manganese substitutions on structural features of parent LaFeO3 and La_{0.7}FeO3 perovskites

The elemental analysis confirms the chemical compositions for the parent LaFeO₃, LaMnO₃ and La_{0.7}FeO₃ perovskites as well as for the La_{0.7}Fe_{0.8}Mn_{0.2}O₃ and La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O₃ perovskites (see Table S2 in SI).

3.1.1. XRD analysis

Fig. 1 shows X-ray diffractograms recorded on single and dual substituted stoichiometric and La-deficient parent perovskites. An orthorhombic perovskite structure is usually characterized with the space group Pnma (PDF 37-1493). The lack of detection of extra reflections emphasizes the absence of significant phases segregations. Minor shift in 2θ values is observed on the most intense (002) reflection on XRD pattern recorded on La_{0.7}Fe_{0.8}Mn_{0.2}O₃ consistent with a slight contraction effect of the unit cell as reported in Table 1. In principle, the substitution of Fe³⁺ by Mn³⁺ should not induce discernible changes on the unit cell because they have comparable ionic radii. By contrast, such effect could be assigned to partial stabilization of Fe⁴⁺ and/or Mn⁴⁺ characterized by lower ionic radii of respectively 0.585 and 0.56 Å vs. 0.64 Å for Fe³⁺ and 0.66 Å for Mn³⁺.

The absence of MnO_x and CaO bulk detectable impurities is also remarkable on dual substituted perovskites (see Fig. 1(B)) emphasizing the fact that Ca²⁺ and Mn³⁺ would be predominantly stabilized in the perovskite lattice. Slight contraction effects are noticeable on La-deficient La_{0.7}Ca_xFe_{0.8}Mn_{0.2}O₃ compositions which accentuate on the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ composition (see Table 1) underlining sharp structural deformations on the stoichiometric composition. Although the topology of substituted perovskites is thermodynamically stable, the structure may distort as a result of preferential bonding of A and B cations [20,21]. Typically, the formation of π (B-O) in A_yA_{1-y}B_xB_{1-x}O₃ can compete with σ (A-O). This competition may also concern σ (B-O) and σ (B'-O) bond formation which means that the covalency of the B-O bond will depend on the nature of the B' and A cations

[22,23]. In general, BO₆ octahedra tilting, bond length change, cationic displacements and anionic vacancies are the main parameters which govern structural deformation of substituted perovskites. The substitution of La³⁺ by Ca²⁺ creates electronic imbalances, while the comparison of their ionic radii (1.36 Å vs. 1.34 Å respectively) should not induce structural changes. Based on this, charge imbalances should be compensated through the creation of anionic vacancies and/or through changes in the valence state of B cation. As aforementioned, the electroneutrality could induce the stabilization of Fe⁴⁺ and/or Mn⁴⁺ and would reasonably explain the contraction effects observed on La-deficient compositions and more importantly on the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ composition.

Table 1. Changes in structural features from XRD measurements induced by single and dual substitutions of parent stoichiometric LaFeO₃ and La-deficient La_{0.7}FeO₃ perovskites.

Catalysts	Crystal system	a(Å) ^a	b(Å) ^a	$c(A)^a$	unit cell volume(Å ³) ^{<i>a</i>}	Crystallite size $(nm)^{b}$
LaFeO ₃	orthorhombic	5.560(1)	7.861(0)	5.557(1)	242.88±0.05	29.4
La _{0.7} FeO ₃	orthorhombic	5.560(0)	7.878(1)	5.535(1)	242.44±0.03	21.7
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	orthorhombic	5.547(0)	7.815(0)	5.536(1)	240.07±0.01	20.4
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	orthorhombic	5.532(1)	7.801(0)	5.519(0)	238.22±0.06	24.8
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	orthorhombic	5.540(0)	7.773(1)	5.513(0)	237.46±0.03	23.7
La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}	orthorhombic	5.491(0)	7.764(1)	5.483(0)	233.78±0.05	21.3

^{*a*} lattice parameters calculated from Rietveld refinement method

^b crystallite size calculated by the Williamson-Hall plot method using LaB₆ as standard specimen



Fig. 1. XRD diffractograms of LaFeO₃, La-deficient $La_{0.7}Fe_{1-y}Mn_yO_3$ perovskites with y = 0 or 0.2 (A), and dual substituted $La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O_3$ perovskites with x = 0.1, 0.2 or 0.4 (B).

3.1.2. Mössbauer spectroscopy

Mössbauer spectra, recorded at room temperature on parent and doped perovskites, are reported in Fig. S2 in SI. Spectra recorded on parent LaFeO₃ and La_{0.7}FeO₃ perovskites are featured by a predominant sextet [24,25] characteristic of cationic Fe³⁺ species in octahedral coordination surrounded by six O^{2-} ions whereas La^{3+} ions occupy eight corners of the orthorhombic unit cell along with a less intense paramagnetic singlet. Distortion of BO₆ octahedra from the ideal cubic structure was confirmed by the quadruple splitting. A visible broadening of line width appears for the Mn-doped perovskites along with marked quadruple splitting measuring the electrical field gradient at the iron site. Such phenomenon is ascribed to local environment effects that lead to anisotropic deformations of the environment of the Fe (+III) species in distorted octahedral sites [26]. Data fitting consisted in the superimposition of several sextets assigned to specific locations of cationic iron species. The contribution of quadrupole-split doublet, with the hyperfine parameters typical of superparamagnetic magnetite particles has been taken into account for manganese substituted perovskite compositions [27,28]. The estimates of the Isomer Shift (IS), the Quadrupole Splitting (QS) and the Hyperfine magnetic Field (HF), fitted from Mössbauer spectra in Fig. S2 in SI, are collected in Table 2. They inform on the nature and coordination geometry of cationic iron species, electronic density and magnetic state. Their relative abundances have been calculated according to a binomial distribution [29]. For Mn-substituted perovskites, the component with high hyperfine field values has been assigned to Fe^{3+} involved in the electron delocalization process in the absence of manganese as nearest neighbors whereas lower HF values correspond to iron species surrounded by manganese. As a general trend, iron preferentially occupies octahedral sites. Nevertheless, significant deviations are remarkable on the distribution between A- and B-sites on La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O₃ characterized by balanced occupancy ratio of octahedral and tetrahedral sites. Battle et al. [30] reported in orthorhombic type perovskite structure that the transition from cubic to orthorhombic structure results from an ordered arrangement of oxygen vacancies along [101] axes with layers of iron cations in tetrahedral coordination. Mere A site deficiency did not incur unusual oxidation state of iron species, indicating a charge compensation likely governed by the formation of oxygen vacancies. The presence of high spin Fe⁴⁺ species seems more related to incorporation of A site cations with lower valence regardless of stoichiometry or deficiency in A site. Isomer shift (IS) values which are characteristic of Fe (+III) species located in the octahedron sites increased with the amount of calcium doping implying that local distortion of octahedron is associated closely with cation doping in A site. This seems in agreement with XRD results that more calcium-doping causes remarkable asymmetry of the crystal structure especially for the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ composition.

Catalyst	Iron species	Coordination geometry	Phase/iron site ^a	Valency	HF (T) ± 0.1T	IS (mm/s) ± 0.05 mm/s	QS (mm/s) ± 0.05 mm/s	Relative abundance (%)
				2.8	55.0	0.57	-0.41	4.83
	Ea ³⁺	Undistorted		3.29	51.9	0.37	-0.04	61.71
Catalyst LaFeO3 La0,7FeO3 La0,7FeO3 La0,7FeO3 La0,6Ca0,1FeO3Mn02O3	ге	octahedron		3.35	48.1	0.32	-0.33	5.34
LaFeO ₃				2.7	47.7	0.58	0	6.68
	Fe ³⁺	Distorted octahedron		3.4	-	0.30	0.38	11.35
	$\mathrm{Fe}^{3+\delta}$	Tetrahedron		3.8	-	0.14	0.16	5.26
	Fe ⁴⁺	High spin		4	37.0	0.07	QS (mm/s) ± 0.05 mm/s a -0.41 -0.04 -0.033 0 0 38 0.16 -0.32 -0.01 -0.02 0.13 0.02 0.02 0.07 -0.02 0.38 -0.03 -0.02 0.78 0.32 0.38 -0.03 -0.02 0.78 0.32 0.38 -0.03 -0.02 0.78 0.32 0.38 -0.03 -0.02 0.02 0.02 0.02 0.10 -0.05 0.11 -0.05 0.11 -0.02 0.577 0.33 0.24 -0.03 0.00 -0.01 -0.02 0.17 -0.09 0.12 0.11 -0.15 0.84 0.24 0.00 0.00	4.83
				3.20	51.6	0.37	-0.01	52.35
	Fa ³⁺	Undistorted		3.30	49.8	0.34	-0.02	13.87
	re	octahedron		3.20	48.7	0.38	0.13	4.10
				3.00	46.3	0.48	0.02	7.40
La _{0.7} FeO ₃				3.05	42.2	0.45	0.07	5.04
240,71 005	2.	Distorted		3.00	18.9	0.48	-0.02	4.78
	Fe ⁵⁺	octahedron		3.10	16.0	0.44	0.02	4.77
				3.00	-	0.47	0.78	3.10
	- 3+ <i>δ</i>	m 1 1		3.30	-	0.35	0.32	2.48
	Fe	Tetrahedron	F 0	3.80	-	0.13	0.38	2.11
	Fe ³⁺	Octahedron	Fe ₂ O ₃ or ferrihydrite		51.1	0.37	-0.03	19.58
	Fe ³⁺	Tetrahedron	(Fe,Mn) ₃ O ₄ -A or		48.1	0.40	-0.02	12.83
Catalyst LaFeO3 La _{0.7} FeO3 La _{0.7} FeO3 La _{0.6} Ca _{0.1} Fe _{0.8} Mn _{0.2} O3	3+ 2+	0.1.1	αFeOOH			0.00	0.02	14.20
	$\frac{\text{Fe}^{3+} \text{ and Fe}^{2+}}{3+}$	Octahedron	(Fe,Mn) ₃ O ₄ -B		46.1	0.39	0.02	14.38
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B		43.5	0.43	0.02	9.60
	Fe^{3+} and Fe^{2+}	Octahedron	(Fe,Mn) ₃ O ₄ -B		42.8	0.23	0.10	7.85
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B		40.0	0.46	-0.06	11.00
	<u>Fe³⁺</u>		Goethite aFeOOH		36.3	0.43	-0.05	9.44
	Fe^{3+} and Fe^{4+}		α (Fe,Mn)OOH		32.9	0.33	0.11	2.31
La _{0.7} Fe _{0.8} Mn _{0.2} O ₃	Fe ³⁺		O-Fe ₂ C or α (Fe,Mn)OOH		15.2	0.30	-0.02	6.04
	Fe ³⁺	Distorted octehedron	γ-FeOOH		-	0.45	0.57	2.81
	Fe ³⁺	Tetrahedron			-	0.19	0.33	3.40
	Fe ³⁺		ferrihydrite		-	0.61	0.24	0.76
	Fe ³⁺		Fe ₂ O ₃ or ferrihydrite		51.4	0.35	-0.03	18.14
	Fa ³⁺	Tetrahedron	Fe ₂ O ₄ -A		49.8	0.34	0.00	14 13
	Fe ³⁺	Tetrahedron	$(Fe,Mn)_3O_4$ -A or		47.6	0.36	-0.01	13.35
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe Mn) ₂ O ₄ -B		45.2	0.43	-0.09	11 59
	Fe^{3+} and Fe^{2+}	Octahedron	$(Fe Mn)_{3}O_{4} B$		42.4	0.18	0.17	8.94
	Fe^{3+} and Fe^{2+}	Octahedron	(Fe Mn) ₂ O ₄ -B		39.3	0.48	-0.09	8.08
La. Ca. Fe. Mr. O.	Fe ³⁺	orunduron	Goethite α FeOOH		36.5	0.32	0.02	4 58
La _{0.6} Ca _{0.1} Fe _{0.8} Mn _{0.2} O ₃	Fa^{3+} and Fa^{4+}		α (Fe Mn)OOH		33.8	0.33	0.11	4 16
	Fe ³⁺		$O-Fe_2C$ or α (Fe Mp)OOH		12.6	0.42	-0.15	5.95
	Fe ³⁺	Distorted	None magnetic		_	0.28	0.84	5.15
	E. ³⁺	octanedron	oxide farribudrita			0.61	0.24	0.64
	Fe Fe ³⁺	Distorted	leitinyurite		-	0.42	0.24	1.30
	P 4+	octahedron	Lish spin			0.05	0.00	2.00
	ге		rugn spin		-	0.05	0.00	3.99

Table 2. Hyperfine parameters from Mössbauer spectroscopy analysis of dual substituted LaFeO₃ and La-deficient La_{0.7}FeO₃ perovskites

	Fe ³⁺	Tetrahedron	Fe ₃ O ₄ -A	49.9	0.25	-0.06	9.29
	Fe ³⁺	Tetrahedron	$(Fe,Mn)_3O_4$ -A or α FeOOH	49.0	0.48	0.03	18.95
	Fe ³⁺	Tetrahedron	$(Fe,Mn)_3O_4$ -A or α FeOOH	48.6	0.17	-0.06	12.87
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	45.1	0.42	-0.04	15.24
Lao ₄ Cao ₂ Feo ₈ Mno ₂ O ₂	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	42.1	0.27	0.09	13.00
240.0040.21 00.811110.203	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	39.1	0.42	-0.06	9.20
	Fe ³⁺		Goethite α FeOOH	34.6	0.58	0.00	6.43
	Fe ³⁺ and Fe ⁴⁺		α (Fe,Mn)OOH	34.4	-0.19	0.26	3.65
	Fe ³⁺	Distorted octahedron	Fe ³⁺ none metallic oxide	-	0.41	0.93	4.19
	Fe ⁴⁺		Fe ⁴⁺ hsp	-	-0.01	0.72	3.35
	Fe ³⁺	Distorted octahedron	γ -FeOOH or Fe ³⁺	-	0.35	0.59	3.83
	Fe ³⁺	Tetrahedron	Fe ₃ O ₄ -A	49.6	0.34	-0.04	8.65
	Fe ³⁺	Tetrahedron	$(Fe,Mn)_{3}O_{4}-A$ or α FeOOH	47.0	0.34	-0.03	16.29
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	44.3	0.33	-0.01	14.28
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	41.9	0.36	-0.03	13.52
	Fe ³⁺ and Fe ²⁺	Octahedron	(Fe,Mn) ₃ O ₄ -B	39.6	0.34	-0.01	11.11
La Ca Ea Ma O	Fe ³⁺		Goethite α FeOOH	36.3	0.42	-0.04	11.18
La _{0.6} Ca _{0.4} Fe _{0.8} IvIII _{0.2} O ₃	Fe ³⁺ and Fe ⁴⁺		α (Fe,Mn)OOH	32.6	0.39	-0.07	8.34
	Fe ³⁺		O-Fe ₂ C ou α (Fe,Mn)OOH	14.2	0.38	0.08	8.74
	Fe ³⁺	Tetrahedron		-	0.26	0.76	1.24
	Fe ³⁺	Tetrahedron		-	0.25	0.31	3.25
	Fe ³⁺		ferrihydrite	-	0.61	0.24	0.68
	Fe ⁴⁺			-	0.09	0.24	2.72

^a -A and -B correspond to two distinct coordination sites for iron in magnetite

3.1.3. Scanning Electron Microscopy (SEM) coupled with Energy-dispersive X-ray spectroscopy (EDS).

SEM images recorded on single and dual substituted stoichiometric and La-deficient perovskite materials are illustrated in Fig. S3 in SI. Fig. 2 reports SEM analysis on La_{0.7}Fe_{0.8}Mn_{0.2}O₃ as example. Similar observations appear whatever the elemental composition. Large aggregates as well as absence of morphological organization is noticeable. Elemental compositions on the volume analyzed from Dispersive X-ray Spectroscopy analysis (1 μ m³) are reported in Table 3. The comparison of the atomic compositions thus obtained with those obtained from ICP-AES analysis (see Table S2 in SI) does not underline significant deviations. This comparison seems to indicate a random distribution of the elements in the volume analyzed. Back scattered electron images reveal no obvious disparity

for calcium and manganese. The same trend is observed on stoichiometric LaFeO₃, La_{0.7}FeO₃ and LaMnO₃ perovskites (Fig. S3(A)-(C) in SI). The absence of large spots that would reflect the formation of large aggregates irrespective of the composition agree with a preferential formation of a single perovskite phase. This confirms a predominant substitution of lanthanum by calcium and iron by manganese inside the perovskite lattice rather than formation of extra-lattice single metal oxides dispersed on the perovskite substrate. All these observations match with those obtained from XRD analyses which did not indicate discernible bulk impurities formation ascribed to manganese oxide segregation.



Fig. 2. SEM image and elemental mapping on La_{0.7}Fe_{0.8}Mn_{0.2}O₃ composition.

catalvet	At	tomic compo	sition (at.	Atomic ratio ^{<i>a</i>}			
Catalyst	La	Ca	Fe	Mn	Mn/Fe	A/B	
LaFeO ₃	48.6	-	51.4	-		0.94 (1.02)	
La _{0.7} FeO ₃	38.0	-	62.0	-		0.61 (0.67)	
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	37.2	-	49.3	13.5	0.40 (0.40)	0.59 (0.71)	
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	31.1	6.4	48.5	13.9	0.29 (0.25)	0.60 (0.68)	
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	29.6	11.3	46.0	13.1	0.28 (0.15)	0.69 (0.78)	
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}\\$	26.3	20.9	40.9	11.9	0.29 (0.25)	0.90 (1.00)	

Table 3. Atomic composition from EDS analysis perovskite samples.

^a calculated from ICP-AES in bracket

3.2. Reducibility and oxygen mobility

3.2.1. Hydrogen Temperature-Programmed Reduction (H2-TPR) experiments

First experiments on benchmark LaFeO₃, La_{0.7}FeO₃ and LaMnO₃ (see Fig. 3(A)) revealed for LaFeO₃ a weak H₂ consumption below 700°C emphasizing unreducible sample in this temperature range. By contrast, two distinct H₂ consumptions on La_{0.7}FeO₃ take place in the temperature ranges 150-425°C and 425-625°C assigned to a two steps reduction of extraframework hematite species: Fe³⁺ \rightarrow Fe²⁺ and Fe²⁺ \rightarrow Fe⁰. For the benchmark LaMnO₃, the H/Mn ratio agrees with the stabilization of Mn⁴⁺ characterized by a two steps reduction process of Mn⁴⁺ to Mn³⁺ and Mn³⁺ to Mn²⁺ respectively in the temperature ranges 250-500°C and above 650°C. This result emphasizes on this sample a current over-stoichiometry of oxygen which agrees with a classical formulae LaMnO_{3+δ}. These preliminary experiments on benchmark catalysts reveal significant overlapping between the reduction of oxidic manganese and iron species which unfortunately cannot lead to accurate calculations neither on the atomic H/Mn and H/Fe ratios nor on the over-stoichiometry of oxygen. Hence, the estimates of the overall H/Mn+Fe have been calculated and the corresponding values are reported in Table 4. The reduction of extra-framework hematite species proceeds on $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ similarly to $La_{0.7}FeO_3$. A discernible broad and weak shoulder arises above 600°C which reflects the reduction of Mn^{3+} to Mn^{2+} . On the basis of this information, the substitution of Fe³⁺ by Mn^{3+} in $La_{0.7}FeO_3$ did not suppress the extraction of iron lattice species but could either slower this process or weaken the stabilization of Mn^{4+} then explaining a lower H/Mn+Fe ratio.

Modifications of redox properties become more apparent when calcium and manganese codoping are implemented, featured by the broadening of reduction peaks and more distinct separation of the reduction processes as exemplified in Fig. 3(B). Indeed, the reduction of Mn^{4+} to Mn^{3+} shifts towards lower temperature while reduction of Mn^{3+} to Mn^{2+} is delayed as calcium doping content increases. The total hydrogen consumption is boosted on $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ (1.44 mmol/g) becoming comparable to that measured on $La_{0.7}FeO_3$ (1.45 mmol/g). At a first glance, it could be seen that $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ mimics the behavior of $La_{0.7}FeO_3$. Nevertheless, a rise in calcium substitution induces lower hydrogen consumption which suggests that calcium substitution would block the extraction/segregation of hematite. Jointly a H₂ consumption process becomes distinguishable above 700°C which intensifies on the stoichiometric $La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_3$ composition emphasizing globally a loss of reducibility assigned either to a lower formation of hematite or a lower stabilization of Mn^{4+} .



Fig. 3. H₂ consumption curves plotted against temperature on parent stoichiometric LaFeO₃ and LaMnO₃ compositions and La_{0.7}Fe_{1-y}Mn_yO₃ with y = 0 and 0.2 (A) and La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O₃ with x = 0.1, 0.2 and 0.4 (B).

3.2.2. Oxygen Temperature Programmed Desorption experiments (O₂-TPD)

O₂-TPD experiments is useful to investigate surface and bulk oxygen mobility. In case of perovskite type materials, two different desorption processes currently appear at low and high

temperature assigned to suprafacial oxygen (α -oxygen) and intrafacial oxygen (β -oxygen) respectively. It was found that α -oxygen desorption would mainly depend on nonstoichiometry and structural defects according to the nature of the A-cations and would correspond to weakly bonded oxygen species [30]. On the other hand, intrafacial oxygen (β oxygen), desorbing at higher temperature, would be more related to the nature of the B cation and usually ascribed to oxygen lattice species, i.e. O^{2-} . As recently reported [11], α -O₂ desorption can overlap β -O₂ desorption process [31,32] which makes uneasy further assignments and quantification. By way of illustration, Fig. 4 shows a broad α -O₂ desorption on LaMnO₃ in the temperature 525-700°C and a faster β -O₂ desorption above 700°C. The former process on La_{0.7}FeO₃ is delayed taking place above 650°C and a second desorption process takes place at higher temperature but further assignment can be unclear because of the coexistence of extra-framework iron oxide species. The broad peak appearing more distinctly on La_{0.7}FeO₃ below 200°C would correspond to the desorption of weakly bonded hydroxyl species and/or physisorbed oxygen species [33]. This process disappears on single and dual substituted samples. This low temperature desorption process has not been taken into account on the calculation of the total amount of desorbed oxygen. The amount of α -O₂ and β -O₂ released during O₂-TPD has been roughly estimated from the decomposition of the overall signals. Results are reported in Table 4. They essentially reveal a predominant β -O₂ desorption.

The substitution of iron by manganese in La_{0.7}FeO₃ leads a much slower desorption of suprafacial oxygen species and a shift to higher temperature of the β -O₂ desorption. Nevertheless, a sharp increase in the amount of oxygen released from the surface is noticeable. Improved bulk oxygen mobility can be rationally explained by the creation of more oxygen vacancies deriving from partial replacement of iron cations by manganese cations. Mössbauer spectroscopic did not reveal the presence of Fe⁴⁺ induced by A site deficiency suggesting a charge compensation mostly governed through change in the oxidation state of manganese and/or formation of oxygen vacancies. A slight contraction effect on the cell volume of La_{0.7}Fe_{0.8}Mn_{0.2}O₃ would agree with the stabilization of Mn⁴⁺ exhibiting a lower ionic ratio than Mn³⁺. Such evolution seems consistent with the desorption features observed on this sample. On the other hand, Fig. 5, illustrating calcium substitution, leads to intriguing observations first related to similar desorption features on La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O₃ and on La_{0.7}FeO₃. An increase in calcium substitution to *x* = 0.2 leads to improved oxygen mobility. The higher amount of desorbed oxygen is consistent with structural changes by considering that more extensive calcium doping should stabilize either iron or manganese cations in B site in higher oxidation state inducing a weakening of the B-O bond. By contrast, the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ behaves differently with a sharp loss of oxygen mobility reaching a minimum value of 0.17 mmol/g and seems in reasonable agreement with a loss of reducibility. Hence, despite the beneficial effect of calcium to changes in the valence state of B-cation, the prevalent parameter in determining the extent of oxygen mobility lies in the A-deficiency.

Quantitative analysis of O_2 -TPD measurements summarized in Table 4 does not indicate that the specific surface area governs the release of oxygen from the surface. Such observation seems in agreement with the prevalence of an intrafacial oxygen desorption at the expense of suprafacial oxygen desorption.



Fig. 4. O_2 -TPD profiles recorded on LaFeO₃ (a); La_{0.7}FeO₃ (b); LaMnO₃ (c); La_{0.7}Fe_{0.8}Mn_{0.2}O₃ (d).

Fig. 5. O₂-TPD profiles recorded on $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ (a) and $La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O_3$ with x = 0.1(b); x = 0.2 (c); x = 0.4 (d)

	eo3 perovskite	·		S _{th} /SSA		H ₂ -TPR ana	alysis	O ₂ desorbed (mmol/g)		
Catalyst composition	Crystallite size d (nm) ^a	SSA (m²/g)	$\frac{S_{th}}{(m^2/g)^b}$		Pore volume (cm ³ /g)	H_2 consumption $(mmol \ ^{g-l})^c$	H/Fe+Mn	α -O ₂ desorption	β -O ₂ desorption	
LaFeO ₃	29.4	15.5	30.8	1.99	0.10	-	-	< 0.01	0.03	
La _{0.7} FeO ₃	21.7	35.5	41.7	1.17	0.10	1.45	0.58	0.03	0.09	
LaMnO ₃	38.7	17.8	23.4	1.31	0.09	2.58	1.25	0.05	0.18	
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	20.4	31.5	44.4	1.41	0.09	1.16	0.47	0.01	0.23	
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	24.8	32.5	36.5	1.12	0.14	1.44	0.55	0.06	0.14	
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	23.7	32.6	38.2	1.17	0.13	1.15	0.45	0.04	0.24	
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}\\$	21.3	31.8	42.5	1.34	0.17	0.77	0.31	0.01	0.16	

Table 4. Textural properties, reducibility and oxygen desorption features on single and dual substituted parent stoichiometric LaFeO₃ and La-deficient La_{0.7}FeO₃ perovskite

^{*a*} crystallite size from XRD measurements - ^{*b*} S_{th} = $6 \times 10^3 / \rho_{\text{perov.}} d$

^{*c*} total H₂ consumption the temperature range 150-700 °C

3.3. Surface properties

3.3.1. Textural properties from nitrogen physisorption

Adsorption isotherms and pore size distribution curves are shown in Fig. S4 in SI. The experimental isotherms of lanthanum ferrite doped with calcium and manganese exhibit a combination of type III and type V isotherms characterized by the absence of B-point at low relative pressure zone yet the presence of hysteresis loop at higher relative pressure zone. The former resulted from the weak interaction between adsorbate and perovskite adsorbent while the latter originated from capillary condensation effect. In all cases, a weak porosity is observable characterized by low pore volume.

Data in Table 4 indicate moderate variations of the specific surface area (SSA) of perovskites doped with manganese in the range of 15.5-35.5 m²/g. Enlarged SSA combined with decreased crystallites size (see Table 4) manifest improved surface properties. A simple model for depicting the morphology of perovskites and the formation of agglomerates can be used. Such information can be useful to explain their ability to store oxygen. It was earlier explained that oxygen availability can be related to a faster diffusion along the grain boundaries. A rough estimation of theoretical surface S_{th} from the crystallites size can be compared to experimental specific surface area (SSA). As reported elsewhere, the S_{th}-to-SSA ratio can underline the fraction of surface lost by agglomeration between crystal domains as earlier explained [34]. As observed in Table 4, no significant deviation is discernible which emphasizes the fact that changes in SSA are essentially due to the growth of the perovskite crystallites.

3.3.2. Surface composition from XPS analysis

The characteristic La 3d, Fe 2p, Mn 2p and Ca 2p core levels have been examined. The corresponding photopeaks are collected in Figs. S5 and S6 in SI respectively on parent samples, on $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ and on the series $La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O_3$. Binding energy (BE) near 834 eV on the La 3d photopeak characterizes trivalent cation La^{3+} stabilized inside the perovskite structure. For Ca 2p core level, BE values agree with the stabilization of Ca^{2+} .

The Fe $2p_{3/2}$ photopeak near 710.2 eV, accompanied by a pronounced satellite structure in the range 718-719 eV, is characteristic of Fe³⁺ cations. Mn $2p_{3/2}$ photopeaks exhibit two components assigned to Mn³⁺ (BE = 641.2 eV) and Mn⁴⁺ (BE = 642.5 eV). [35,36]. Mn 3s photopeak in Fig. S7 in SI provides useful insights on the oxidation state of manganese. Multiplet splitting of Mn 3s peak results from the coupling of non-ionized Mn 3s electrons and 3d valence band electrons [37]. In theory, multiplet splitting can be observed for Mn²⁺, Mn³⁺ and Mn⁴⁺ while Mn⁵⁺ is featured by less defined splitting or broadening. The average oxidation state (AOS) of manganese in the perovskite has been estimated according Eq. (1).

Average Oxidation State (AOS) =
$$8.956 - 1.12\Delta E_{\text{splitting}}$$
 (Mn 3s) (1)

The variation of Mn AOS in perovskite has been closely associated to the nature of A-site and stoichiometry. In general, La-deficiency favors an increase of Mn AOS. To check this assertion spectral features reported on LaFe_{0.8}Mn_{0.2}O₃ have been included in Table 5 for further comparison with La_{0.7}Fe_{0.8}Mn_{0.2}O₃. This seems in agreement with H₂-TPR showing that La-deficiency can improve the reducibility of the perovskite. Calcium doping with x =0.1 increases Mn AOS which is supposed to be assigned to the synergistic effect of calcium and manganese. As a matter of fact, the average oxidation state (AOS) of manganese cations for the series La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O₃ is mainly determined by the overall cationic charge in A site and continuously decreases reaching a minimum for La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃. It is thereby assumed that the reducibility of manganese species located in the lattice is affected primarily by the composition of A site. This is supported by a linear increase of the Mn AOS as a function of the total H_2 uptake from H_2 -TPR as exemplified in Fig. 6.

Comparison between the O 1s photopeak and the amount of O₂ released from O₂-TPD experiments is not obvious likely due to the predominance of intrafacial β -O₂ desorption. O 1s photopeak is usually characterized by two components ascribed to lattice oxygen species near 529 eV (O_{lattice}) and surface O-containing species near 531 eV (O_{ads}). The highest value obtained on La_{0.7}FeO₃ is consistent with a low temperature desorption below 250°C. On the other hand, no clear comparison appears with respect to Mn and Ca substitution revealing some discrepancies between the evolution of the values for O_{ads}-to-O_{lattice} ratio with respect to the amount of oxygen released.

Finally, no significant deviation appears by comparing the surface atomic Mn/Fe ratios with those deduce from elemental analysis (see Table S2 in SI) which seems to indicate the absence of preferential segregations of Mn and Fe. Regarding the A/B ratio, some deviations appear distinctly on stoichiometric compositions emphasizing a significant lanthanum surface enrichment.

	Binding energy (eV)								Semi-quantitative analysis				
Catalyst	La 3d	Ca 2p	Fe 2p	Mn 2p	$\Delta E_{splitting}\left(eV\right)$	Mn AOS	O 1s (O _{ads})	O 1s (O _{lattice})	Ca/La	Mn/Fe	A/B	O _{ads} /O _{lattice}	
LaFeO ₃	833.8	-	710.2	-	-	-	531.1	529.2	-	-	1.34 (1.02)	1.28	
La _{0.7} FeO ₃	834.2	-	710.9	-	-	-	531.2	529.7	-	-	0.48 (0.67)	1.61	
$LaFe_{0.8}Mn_{0.2}O_3$	8.34.0	-	710.3	641.7	5.31	2.98	531.5	529.6	-	0.38(0.25)	1.85(0.99)	0.80	
$La_{0.7}Fe_{0.8}Mn_{0.2}O_{3}\\$	834.1	-	710.5	641.7	5.12	3.19	531.3	529.7	-	0.32 (0.25)	0.67 (0.71)	0.76	
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	833.9	346.7	710.6	641.8	4.95	3.38	531.4	529.6	0.18 (0.17)	0.32 (0.25)	0.64 (0.68)	0.89	
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	833.7	346.4	710.2	641.7	5.28	3.01	531.3	529.4	0.31 (0.34)	0.43 (0.25)	0.74 (0.78)	1.10	
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}$	833.6	346.4	710.2	641.8	5.62	2.63	531.3	529.0	0.68 (0.38)	0.41 (0.25)	1.27 (1.00)	0.75	

Table 5. XPS analysis of single and dual substituted calcium manganese LaFeO₃ and La_{0.7}FeO₃ perovskites

Elemental composition in bracket



Fig. 6. Comparison between the reducibility and the average oxidation state of manganese (Mn AOS).

3.4.1. Oxidative properties

Four successive temperature programmed reaction (TPR) experiments were performed in stoichiometric (STO1), lean, rich and stoichiometric (STO2). Prior TPR experiments, the samples were pre-activated in air at 600°C. Conversion profiles vs. temperature for CO, propene and propane oxidation are reported in Fig. 7. STO1 and STO12 profiles are comparable for CO and propene oxidation underlining the absence of significant deactivation due to surface changes in terms of composition and valence state of active elements. Ca- and Mn-substitutions induce significant gain in conversion compared to the benchmark La_{0.7}FeO₃ and LaFeO₃ catalysts. La-deficiency has a beneficial effect as well as Mn-substitution. On the other hand, Ca-substitution has no discernible effect for CO oxidation compared to La_{0.7}Fe_{0.8}Mn_{0.2}O₃ and a slight detrimental effect is even observed on the rate of propene conversion recorded on the series La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O₃. The light-off temperatures collected in Table 6 provide a qualitative overview of the catalytic performances highlighting the superiority of La_{0.7}Fe_{0.8}Mn_{0.2}O₃.

			-	-							
Catalyst	Light-off temperature T_{50} (°C)										
Catalyst		STO1			STO2						
	СО	C_3H_6	C_3H_8	СО	C_3H_6	C_3H_8					
LaFeO ₃	480	477	500	487	475	488					
La _{0.7} FeO ₃	395	448	>520	399	456	483					
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	360	405	518	359	415	514					
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	366	413	513	369	441	503					
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{.2}O_{3}$	375	471	513	388	475	510					
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}\\$	375	462	479	366	458	490					

Table 6. Light-off temperature (T_{50}) of iron-based perovskite samples.



Fig. 7. Catalytic performances of single and dual substituted iron-based perovskites in oxidation reactions : \bullet LaFeO₃; \bullet La_{0.7}FeO₃; \checkmark La_{0.7}FeO₈Mn_{0.2}O₃; \diamond La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O₃; \diamond La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O₃; \diamond La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃.

Kinetics was analyzed based on Temperature-Programmed CO and propene conversions recorded in stoichiometric conditions at 320°C and 375°C respectively. As described in SI, the mass balance equation of a plug flow reactor and a first order kinetics [38-40] have been assumed for the calculation of the specific and normalized rates, respectively labeled $r_{spec.}$ and $r_{norm.}$ The boundary conditions given by the Weiss-Prater criterion according to Eq. (S4) in SI

was also verified showing the absence of significant perturbations induced by mass transfer phenomena.

Clearly, the evolutions strictly agree with previous tendencies reflecting the highest catalytic properties of $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ and a detrimental effect of calcium on the reaction rates more accentuated in propene oxidation. The comparison of the pre-exponential factor (A) and apparent activation energy (E_{app}) values of the rate constant k for CO and propene oxidation explain this rate enhancement by a lowering of E_{app} . The pre-exponential factor values in Tables 7 and 8 seem to be independent of the specific surface area and mostly depend on the elemental composition.

			STO1		STO2						
Catalyst	$\begin{array}{c} S_{BET} \\ (m^2g^{-1}) \end{array}$	$r_{spec.} a (mol s^{-1} g^{-1})$	Normalized rate ^{<i>a</i>} $r_{norm.}$ (mol s ⁻¹ m ⁻²)	A^{b} (m ³ s ⁻¹ g ⁻¹)	E _{app} ^b (kJ mol ⁻¹)	$r_{\text{spec.}}^{a}$ (mol s ⁻¹ g ⁻¹)	Normalized rate ^{<i>a</i>} $r_{norm.}$ (mol s ⁻¹ m ⁻²)	A^{b} (m ³ s ⁻¹ g ⁻¹)	E _{app} ^b (kJ mol ⁻¹)		
LaFeO ₃	15.5	0.1×10 ⁻⁷	0.6×10 ⁻⁹	3.6×10 ³	121.4	0.2×10 ⁻⁷	0.8×10 ⁻⁹	2.9×10 ³	162.4		
La _{0.7} FeO ₃	35.5	3.4×10 ⁻⁷	9.4×10 ⁻⁹	3.3×10 ³	107.9	1.8×10 ⁻⁷	5.2×10 ⁻⁹	43.0×10 ³	123.3		
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	31.5	7.8×10 ⁻⁷	24.7×10 ⁻⁹	2.0×10 ³	99.1	7.8×10 ⁻⁷	24.7×10 ⁻⁹	3.6×10 ³	102.4		
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.5	7.2×10 ⁻⁷	22.3×10-9	3.0×10 ³	102.2	2.7×10 ⁻⁷	8.3×10 ⁻⁹	3.2×10 ³	104.3		
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.6	5.6×10-7	17.0×10 ⁻⁹	2.2×10 ³	102.1	4.0×10 ⁻⁷	12.2×10 ⁻⁹	3.3×10 ³	106.6		
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}$	31.8	5.5×10 ⁻⁷	17.2×10 ⁻⁹	8.2×10 ³	108.9	5.6×10 ⁻⁷	17.5×10 ⁻⁹	2.9×10 ³	102.6		

Table 7. Impact of single and dual substitution on the kinetic features of LaFeO₃ and La_{0.7}FeO₃ in CO oxidation.

^{*a*} calculated at $T = 320^{\circ}C$

^{*b*} $k = A \exp(-E_{app}/RT)$

			STO1		STO2						
Catalyst	S_{BET} (m^2g^{-1})	r _{spec.} ^{<i>a</i>} (mol s-1 g-1)	Normalized rate ^{<i>a</i>} r _{norm.} (mol s ⁻¹ m ⁻²)	A^{b} (m ³ s ⁻¹ g ⁻¹)	E _{app} ^b (kJ mol ⁻¹)	$r_{\text{spec.}}^{a} (\text{mol} \ s^{-1} g^{-1})$	Normalized rate ^{<i>a</i>} r _{norm.} (mol s ⁻¹ m ⁻²)	A ^b (m ³ s ⁻¹ g ⁻¹)	E _{app} ^b (kJ mol ⁻¹)		
LaFeO ₃	15.5	1.1×10 ⁻⁸	6.9×10 ⁻¹⁰	7.1×10 ³	125.0	0.8×10 ⁻⁸	5.3×10 ⁻¹⁰	1.2×10^{4}	129.6		
La _{0.7} FeO ₃	35.5	1.7×10 ⁻⁸	4.88×10 ⁻¹⁰	7.5×10^{3}	122.6	1.3×10 ⁻⁸	3.6×10 ⁻¹⁰	4.5×10^{4}	134.0		
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	31.5	6.7×10 ⁻⁸	21.3×10 ⁻¹⁰	1.8×10 ³	106.7	5.6×10 ⁻⁸	17.6×10 ⁻¹⁰	8.9×10 ³	117.1		
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.5	4.0×10 ⁻⁸	12.2×10 ⁻¹⁰	6.6×10 ⁶	154.6	2.0×10 ⁻⁸	6.3×10 ⁻¹⁰	1.7×10^{5}	139.0		
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.6	1.3×10 ⁻⁸	3.9×10 ⁻¹⁰	6.4×10^{4}	136.0	0.8×10 ⁻⁸	2.4×10 ⁻¹⁰	4.2×10^{3}	123.0		
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}\\$	31.8	1.8×10 ⁻⁸	5.6×10 ⁻¹⁰	1.9×10 ³	115.4	1.0×10 ⁻⁸	3.1×10 ⁻¹⁰	4.1×10 ⁵	145.4		

Table 8. Impact of single and dual substitution on the kinetic features of LaFeO₃ and La_{0.7}FeO₃ in propene oxidation.

^{*a*} calculated at $T = 320^{\circ}C$

^{*b*} $k = A \exp(-E_{app}/RT)$

3.4.2. Reductive properties

NO conversion curves vs. temperature are reported in Fig. 8. NO conversion is weakly detected in lean conditions and slightly improved in stoichiometric conditions on La_{0.7}Fe_{0.8}Mn_{0.2}O₃ and La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O₃. Let us note that for both catalysts, the conversions of propene and CO, in Fig. 7 start nearly at the same temperature than that of NO but conversion curves diverge significantly with a rise in temperature emphasizing the predominant CO and propene oxidation by oxygen. Such observation would agree with a preferential NO reduction by CO at low temperature while at high temperature oxidation reaction by O₂ predominates. By comparing STO1 and STO2, a loss of NO conversion is observed in rich conditions and still emphasizes the superiority of La_{0.7}Fe_{0.8}Mn_{0.2}O₃ compared to La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O₃ in rich conditions while this composition was found inactive in lean and stoichiometric regimes.

By examining the selectivity behavior, no drastic change occurs on $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ and $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$. Both samples exhibit the same selectivity behavior in lean and stoichiometric conditions (see Fig. 9) with a gradual decrease of N₂O production to the benefit of N₂ vs. temperature. Such changes would not be likely related to the successive N₂O reduction. Indeed, this sub-reaction becomes usually predominant at high NO conversion and would be consequently slower in the residual conversions (being lower than 10%) recorded in our operating conditions. Accordingly, the predominant production of N₂ with a rise in temperature would agree preferentially with a higher activation barrier than that of N₂O formation. In rich conditions, the predominant production of ammonia takes place at high temperature and agrees with fast NO dissociation and simultaneous hydrogenation of adsorbed N atoms by hydrogen instead of N₂ production through the recombination of two

chemisorbed nitrogen atoms. The production of ammonia can be reasonably explained if a source of hydrogen is available. An extra production of H_2 can be originated from reforming or partial oxidation of hydrocarbons in rich conditions. Such hypothesis is supported by the examination of CO conversion profiles vs. temperature in Fig. 7. Indeed, reforming reactions lead to CO as primary product. Accordingly, the sharp loss in CO conversion observed in rich conditions could be reasonably explained by extra CO production through these reactions. Accordingly, a shift from the CO/NO reaction to the NO/H₂ reaction would occur with a rise in temperature in rich conditions likely due to a predominant partial oxidation of propene to CO and H_2 instead of total oxidation.



Fig. 8. Catalytic performance of single and dual substituted iron-based perovskites in NO reduction reaction: • LaFeO₃; • La_{0.7}FeO₃; • La_{0.7}FeO₃; • La_{0.7}FeO₃; • La_{0.6}Ca_{0.1}FeO₈Mn_{0.2}O₃; • La_{0.6}Ca_{0.1}FeO₈Mn_{0.2}O₃; • La_{0.6}Ca_{0.4}FeO₈Mn_{0.2}O₃.



Fig. 9. Selectivity for $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ (magenta) and $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ (orange) in stoi1, stoi2 and rich conditions. Magenta solid line solid square $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ N₂ selectivity (\blacksquare); Magenta dash line hollow square $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ N₂O selectivity(\square); Magenta dot line big semi-solid square $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity (\blacksquare); Orange solid line solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ N₂ selectivity(\bigcirc); Orange dash line hollow circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc); Orange dot line big semi-solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc); Orange dot line big semi-solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc); Orange dot line big semi-solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc); Orange dot line big semi-solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc); Orange dot line big semi-solid circle $La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_3$ NH₃ selectivity(\bigcirc).

3.4.3. Discussion on structure-reactivity relationship

The establishment of structure-reactivity relationships for perovskite-type materials is never easy because of the versatility of these materials. Most of the examples dealt with catalytic oxidation reactions. Catalytic properties of perovskite-type materials are governed by the presence of defective sites, which ease adsorption, bulk and surface mobility of oxygen, and the stabilization of cations in unusual valence improving their redox properties. Based on this, two distinct reaction mechanisms can be envisioned. The first one involves lattice oxygen periodically restored at the end of continuous cycles, whereas the second one involves adsorbed gaseous oxygen species and depends on the symmetry of atomic orbitals to be conductive for the reaction [41].

The comparisons on benchmark LaFeO₃ and La_{0.7}FeO₃ perovskites reveal the beneficial impact of La-deficiency in propene and CO oxidation reactions. An additional gain in activity is clearly observed after substitution of Fe by Mn in La_{0.7}FeO₃. By contrast, dual substitution does no longer evidence a synergy effect on the activity in CO and propene conversion. On the contrary Lanthanum substitution by Calcium has a slight detrimental effect on the rate of

CO oxidation which becomes more intense for propene oxidation. Similar trends are observable in STO1 and STO2 series. The weak deviations on the rate values recorded during STO1 and STO2 experiments emphasizes stable surface compositions and the absence of significant alterations of the oxidation state of active elements during lean/rich cycles.

Going back to changes on structural and surface properties induced by single and dual substitutions, clearly calcium substitution provokes sharp structural changes related to contraction of the cell volume at increasing Ca concentration. This contraction effect, more significant for the stoichiometric $La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_3$ composition, has not been related to exsolution and segregation processes but mainly to the stabilization of Fe⁴⁺, detected from Mössbauer spectroscopy and characterized by a lower ionic radius than Fe³⁺. The correlative creation of oxygen vacancies to compensate charge imbalance can accentuate this contraction effect. Similarly, the stabilization of Mn⁴⁺ at the expense of Mn³⁺, characterized by a lower ionic radius (0.56 Å vs. 0.66 Å for Mn³⁺) can equally explained these contraction effects. Such hypotheses have been previously envisioned by Onrubia et al. [1] who privileged an increase of the oxidation state of manganese rather than formation of anionic vacancies for explaining the better catalytic properties of Sr-doped perovskites in NO reduction. Clearly, our results differ from those obtained by these authors because La^{3+} substitution by Ca^{2+} leads to lower rates in NO conversion, the most active sample being $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$.

H₂-TPR and O₂-TPD measurements are bulk sensitive characterizing the reducibility and oxygen mobility. No reliable comparison can be drawn with rate values in Table 9. Quantitative comparisons between H₂ uptake from H₂-TPR experiments and the amount of O₂ release from desorption according to the degree of substitution is not obvious. Interestingly, H₂ uptake diminishes at increasing Ca doping which could contradict a greater formation of Mn⁴⁺. In practice we demonstrate that calcium also blocks the formation of more reducible hematite promoted on La-deficient perovskite samples. Regarding O₂-TPD experiments, O₂

desorption is promoted on La-deficient perovskites more prone to form anionic vacancies and occurs more readily on $La_{0.6}Ca_xFe_{0.8}Mn_{0.2}O_3$ with x = 0.1 and 0.2 at lower temperature. On the other hand, opposite trends are observed on the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ composition. Nakamura et. al [42] proposed on $La_{1-x}Sr_xCoO_{3-\delta}$ that increasing the perovskites reducibility decreases its propensity to refill its oxygen vacancies. This suggests that a poorer activity could be expected in case of intrafacial mechanism. Obviously, neither the bulk reducibility of the B cation nor the bulk oxygen mobility would act as critical parameters based on the comparison of H₂ uptake and oxygen release for explaining the impact of single and dual substitutions on the catalytic properties. The absence of correlation between catalytic properties and bulk physicochemical properties has been already pointed out on LaCo_{1-x}Fe_xO₃ [35] for the methane oxidation reaction preferentially obeying to a suprafacial mechanism. Nevertheless, the comparison of rate values with specific surface area is also unable to explain the trends observed of the catalytic properties. Indeed, the rates as well as the pre-exponential factor values of the rate constant vary independently with the specific surface area emphasizing the primacy of the surface composition and oxidation state of active elements. Both parameters would act as critical parameters in determining the catalytst properties. Nevertheless, an important question arises related to the involvement of the mixed oxidation states Fe⁴⁺/Fe³⁺ and Mn⁴⁺/Mn³⁺ in particular the one that will prevail.

Tables 9 compares kinetic data with physicochemical properties. An important information comes from the comparison of $La_{0.7}FeO_3$ and $LaFeO_3$ in CO oxidation which clearly demonstrates that the rate enhancement observed on the former composition cannot be formally explained by the presence of Fe⁴⁺ in the bulk and/or at the surface. Such tendency persists for propene oxidation but in lesser extent. Based on this, any participation of Fe(+IV) as active species can be ruled out in the oxidative properties of Ca-doped samples. Subsequent, rate improvements on $La_{0.7}Fe_{0.8}Mn_{0.2}O_3$ both for oxidation and reduction

reactions emphasize the prevalent role of high oxidation state of manganese corresponding to higher rates and lower activation barriers for CO and propene oxidation. Mössbauer spectroscopy did not detect for this sample the presence of Fe^{4+} which seems in relative agreement with previous investigation reported by Leontiou et al. [11] who observed that the highest activity in NO reduction by CO correspond to the lowest amount of Fe^{4+} and the highest amount of Fe^{3+} .

As found, Ca substitution leads to lower rates explained by an increase in E_{app}. Correlatively, a significant decrease in the manganese average oxidation state is noticeable with a predominant stabilization of Mn(+III). Such a tendency could be responsible for the slight increase observed in Mn/Fe surface composition. Accordingly, а slight aggregation/segregation of small clusters of Mn₂O₃ or MnO cannot be excluded. Alternately, this evolution on the relative surface concentration of manganese could be related to lower extent of hematite segregation. This is due to the stabilizing effect of calcium maintaining iron stabilized as Fe⁴⁺ in octahedral position of the perovskite. Such changes in physicochemical properties could rationally explain the loss of catalytic properties due to calcium substitution especially for the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ compositions.

To summarize, A-site deficiency coupled to manganese doping in B site results in better reducibility of lanthanum ferrite that could create more oxygen vacancies essential for the adsorption of NO and O_2 in competition. In parallel, a cooperation between iron and manganese is suggested for the stabilization of Mn^{4+} at the vicinity of anionic vacancies. Clearly, Ca doping has a detrimental impact because electronic imbalance would be more compensated by an increase of the oxidation state of Fe³⁺ to Fe⁴⁺ as observed from Mössbauer spectroscopy. This could explain the smallest $O_{ads}/O_{lattice}$ ratio for the stoichiometric composition $La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_3$. Hence, low anionic vacancies jointly with low Mn AOS

can reasonably explain the low catalytic performance of this catalyst both for oxidation and reduction reaction.

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	Sper	Unit call			0					STO1			STO2	
Catalysts	$(m^2 g^{-1})$	volume (Å ³)	Fe ⁴⁺ /Fe ³⁺	H ₂ uptake (mmol/g)	desorbed (mmol/g)	Mn AOS	O _{ads} /O _{latt.}	Mn/Fe	$r_{spec.}$ (mol s ⁻¹ g ⁻¹)	$A (m^3 s^{-1} g^{-1})$	E _{app} (kj mol ⁻¹)	$r_{spec.} (mol s^{-1} g^{-1})$	$A (m^3 s^{-1} g^{-1})$	E _{app} (kj mol ⁻¹)
CO oxidation														
LaFeO ₃	15.5	242.88	4.83	0	0.04		1.28		0.2×10 ⁻⁷	3.6×10 ³	121.4	0.2×10 ⁻⁷	2.9×10 ³	162.4
La _{0.7} FeO ₃	35.5	242.44	0	1.45	0.14		1.61		1.8×10 ⁻⁷	3.3×10 ³	107.9	1.8×10 ⁻⁷	43.0×10 ³	123.3
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	31.5	240.07	0	1.16	0.23	3.19	0.76	0.32	7.8×10 ⁻⁷	2.0×10 ³	99.1	7.8×10 ⁻⁷	3.6×10 ³	102.4
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.5	238.22	3.99	1.44	0.23	3.38	0.89	0.32	2.7×10 ⁻⁷	3.0×10 ³	102.2	2.7×10 ⁻⁷	3.2×10^{3}	104.3
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.6	237.46	3.35	1.15	0.33	3.01	1.10	0.43	4.0×10 ⁻⁷	2.2×10 ³	102.1	4.0×10 ⁻⁷	3.3×10 ³	106.6
$La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O_{3}\\$	31.8	233.78	2.72	0.77	0.16	2.63	0.75	0.41	5.6×10 ⁻⁷	8.2×10 ³	108.9	5.6×10 ⁻⁷	2.9×10 ³	102.6
Propene oxidation														
LaFeO ₃	15.5	242.88	4.83	0	0.04		1.28		1.1×10 ⁻⁸	7.1×10 ³	125.0	0.8×10 ⁻⁸	1.2×10 ⁴	129.6
La _{0.7} FeO ₃	35.5	242.44	0	1.45	0.14		1.61		1.7×10 ⁻⁸	7.5×10 ³	122.6	1.3×10 ⁻⁸	4.5×10 ⁴	134.0
$La_{0.7}Fe_{0.8}Mn_{0.2}O_3$	31.5	240.07	0	1.16	0.23	3.19	0.76	0.32	6.7×10 ⁻⁸	1.8×10 ³	106.7	5.6×10 ⁻⁸	8.9×10 ³	117.1
$La_{0.6}Ca_{0.1}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.5	238.22	3.99	1.44	0.23	3.38	0.89	0.32	4.0×10 ⁻⁸	6.6×10^{6}	154.6	2.0×10 ⁻⁸	1.7×10^{5}	139.0
$La_{0.6}Ca_{0.2}Fe_{0.8}Mn_{0.2}O_{3}\\$	32.6	237.46	3.35	1.15	0.33	3.01	1.10	0.43	1.3×10 ⁻⁸	6.4×10 ⁴	136.0	0.8×10 ⁻⁸	4.2×10 ³	123.0
La_0.6Ca_0.4Fe_0.8Mn_0.2O_3	31.8	233.78	2.72	0.77	0.16	2.63	0.75	0.41	1.8×10 ⁻⁸	1.9×10 ³	115.4	1.0×10 ⁻⁸	4.1×10^{5}	145.4

Table 9. Comparison between physicochemical and catalytic properties for CO and propene oxidation

4. Conclusion

Lanthanum ferrite perovskite with orthorhombic structure was prepared by conventional citric method. Such crystal structure was preserved after partial substitution in A- or B-site. Parent LaFeO₃ and La_{0.7}FeO₃ perovskites reveal that La-deficiency promotes the formation of extraframework hematite species. Mn doping in B-site of La-deficient La_{0.7}FeO₃ perovskite did not result in significant phase segregation irrespective to Mn doping degree or A-site composition. By contrast, contraction of unit cell volume and significant structural distortions characterized from Mössbauer spectroscopy appear especially on the stoichiometric La_{0.6}Ca_{0.4}Fe_{0.8}Mn_{0.2}O₃ composition. The stabilization of Fe⁴⁺ proceeds on stoichiometric composition and improves with partial substitution of La by Ca in A-site. Average oxidation state (AOS) of doped Mn species also depends heavily on A-site composition. It is noticeable that Mn⁴⁺/Mn³⁺ ratio changes according to the elemental composition of the solid and that La-deficiency favors an increase of Mn⁴⁺/Mn³⁺ ratio. By contrast, a rise in Ca content decreases the Mn⁴⁺/Mn³⁺. With regard to oxidative properties and reductive properties, La_{0.7}Fe_{0.8}Mn_{0.2}O₃ represents the best formulation with enhanced reaction rates associated with lower apparent activation energy values. Minor deviation between STO1 and STO2 underlines the high stability of the surface properties with no obvious segregation after lean/rich cycle. As general trend, the catalytic activity is not governed by the extent of surface area but mostly by the surface composition, ie. the valence of manganese and the La-deficiency. Subsequent Ca-substitution induces a detrimental effect on the oxidation and reduction reaction rates reasonably explained by a decrease of the oxidation state of manganese. Jointly, the formation of Fe⁴⁺ is

 Mn^{4+}/Mn^{3+} couple than Fe^{4+}/Fe^{3+} couple.

still detected emphasizing the fact that the redox properties are rather governed by the

Despite, the reduction of NO is slow, it clearly appears that the CO/NO reaction predominates at low temperature. On the other hand, the NO/H₂ reaction prevails at high temperature especially in rich conditions when an extra production of hydrogen from partial oxidation and reforming reactions of propene and propane take place predominantly. The involvement of such sub-reactions has a detrimental impact on the selectivity towards N_2 production with a significant production of ammonia.

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Graphical Abstract

