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Screening Divalent Metals for A- and B-Site Dopants in LaFeO₃.

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ABSTRACT: Doping LaFeO₃, a mixed ionic electronic conductor, can serve to increase its ionic and electronic conductivity, as observed in La_{1-x}Sr_xCo_{1-y}Fe_yO_{3-δ} (LSCF), a promising intermediate temperature solid oxide fuel cell (IT-SOFC) cathode material. In this study *ab initio* methods have been employed to assess the viability of a range of divalent A- and B-site dopants for promoting ionic and electronic conductivity, through calculating solution energies and binding energies to charge compensating species. For the A-site, we find that all alkali earth metals considered promote increased conductivity properties, but strontium and calcium have the lowest solution energies and therefore will be suitable dopants, in full agreement with experiment. Surprisingly we find manganese, which has typically been assumed to dope exclusively on the B-site, to have significant probability, based on energetic considerations, to occupy the A-site and be equally as energetically favourable as the traditional strontium dopant under certain conditions. For the B-site, cobalt and nickel, were found to be suitable dopants, promoting ionic and electronic conductivity, due to the variable oxidation state of transition metals. Magnesium also increases conductivity as a B-site dopant in contrast with the other alkali earth dopants studied, which favour the A-site. By considering two compensation mechanisms, O²⁻ vacancy and hole compensation, we show both oxygen vacancies and holes will be promoted in the doped system, in agreement with the experimentally observed mixed ionic electronic conducting properties of doped systems, including LSCF.

1. Introduction

Solid oxide fuel cells (SOFCs) show distinct promise as a clean and renewable alternative to burning fossil fuels. They are efficient, can use a variety of fuels, and have proved to be reliable when operated continuously. (1,2) SOFCs have the standard three main components: a cathode, an electrolyte and an anode. During operation, oxygen is reduced at the cathode site to form O²⁻ ions which migrate through the electrolyte to the anode where they react with a fuel. (1,3) This conduction of ions is a thermally activated process, hence high operating temperatures are necessary, leading to a number of drawbacks including high cost of materials and short lifetimes. The discovery of novel materials for each component is an active field of research, with the aim of bringing operating temperature down to between 500 and 800°C. (2)

As temperatures are reduced, the rate limiting step becomes the reduction of oxygen on the cathode surface; therefore finding cathode materials that catalyse this reaction at lower temperatures is essential to achieve successful intermediate temperature SOFCs (IT-SOFCs). Mixed ionic electronic conductors (MIECs), such as perovskite LaFeO₃, show strong potential as cathode materials as the ionic conductivity serves to increase the area within which the oxygen reduction reaction can take place, increasing the activity of the oxygen reduction reaction at the cathode. In particular, when doped, as in for example La₁-_xSr_xCo_{1-y}Fe_yO_{3-δ} (LSCF), high electronic and ionic conductivities have been observed within the intermediate temperature range, along with a high activity towards oxygen reduction. (4-7) Indeed, LSCF-based cathode materials have demonstrated similar power densities to state-of-the-art La_{1-x}Sr_xMnO_{3-δ} (LSM) for operating temperatures 100°C lower and have a higher tolerance to chromium species, which are often present in common interconnect materials, leading to a more stable performance. (5, 8) However, degradation is still an issue is these materials, (4, 9) although it is proposed that altering their composition could

prevent degradation and improve long term stability. (9) With this in mind, understanding doping in the parent material is of primary importance in order to propose design strategies, an area in which computational studies can provide critical insight. (10-16)

Doping LaFeO₃ affects its structural, transport and electrical properties. (17-30) Doping the A-site with divalent metals, in particular the alkali earths, has been shown to increase the oxygen vacancy concentration, and therefore oxide ion conductivity. (17,18,19) While doping on the B-site, for which the focus is first row transition metals, has been shown to affect magnetic properties and electrical conductivity. (17,20,21,22) However there is no consensus on the most appropriate dopant to use for each site. For example, Ortiz-Vitoriano et al. (23) suggest Ca is the optimum A-site dopant of the alkali earth metals, as it has comparable properties to the strontium doped system while being cheaper, whereas Bidrawn et al. (19) report calcium doped LaFeO₃ to have a lower ionic conductivity compared to strontium doped LaFeO₃ along with a higher activation energy for ionic transport.

Computational studies on doped LaFeO₃ are limited and tend to focus on dopant concentration without first considering the most appropriate dopant for this system. (18, 31) Work that has examined a range of dopants has generally been carried out using interatomic potential based methods. (32) However, using these approaches to model transition metal ions accurately can pose challenges especially when considering both electronic and point defect compensation of the effective charge of the dopant.

In this paper, a range of divalent dopants in LaFeO₃, on the A- and B-site have been investigated using density functional theory (DFT) in order to establish the most appropriate dopant for each site. Manganese, cobalt, nickel and copper are considered for both the A- and B-sites, while magnesium, calcium, strontium, barium and iron are also considered for the A-site;

the only alkali earth metal considered for the B-site is magnesium. Although we would not expect the transition metals to substitute on the A-site of this material, due to their small ionic radii, they have been included in this work for completeness and to investigate the possibility of these impurities occupying both sites when incorporated in LaFeO₃. Solution energies have been calculated, along with the binding energies of these dopants to oxygen vacancies and holes. For the A-site, Sr²⁺ and Mn²⁺ are calculated to be the most energetically favourable from each group of dopants, whereas Co²⁺, Ni²⁺ and Mg²⁺ are the most favourable B-site dopants, from the range of divalent metals studied.

2. Simulation Methods

The ambient temperature structure of LaFeO₃ is the orthorhombic perovskite form, *Pbnm* no. 62,⁽³³⁾ shown in Figure 1, and has been adopted throughout our investigation. Calculations were carried out using *ab initio* based methods, as outlined below.

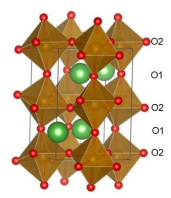


Figure 1. Orthorhombic form of LaFeO₃; lanthanum are shown in green, iron in gold and oxygen in red. The two oxygen environments; O1 and O2 are shown to the right of the structure.

All electronic structure calculations were performed, as in ref. [34], using spin-polarised Kohn-Sham DFT(35,36) and DFT+U⁽³⁷⁾ using the plane-wave pseudopotential technique with the projector augmented-wave⁽³⁸⁾ (PAW) approach to model the core-valence electron interaction, as implemented in the Vienna ab initio simulation package (VASP). (39, 40, 41) The valence configurations used were: La $(5s^25p^66s^25d^1)$, Fe $(4s^23d^8)$, O $(2s^22p^4)$. The electron exchange and correlation was evaluated within the generalised gradient approximation (GGA+U) using the functional of Perdew, Burke and Ernzerhof (PBE), (42) which was found to reproduce the experimental structure of LaFeO₃ most accurately. (34) A k-mesh of (4×4×3) generated by the Monkhorst-Pack(43) scheme is used for Brillouin zone sampling for the LaFeO₃ unit cell. Integration over the first Brillouin-zone used Gaussian smearing ($\sigma = 0.05$) during structural relaxations. The plane-wave basis set was converged at a kinetic cutoff of 650 eV, with the total energy converging to within 0.001 eV per atom. A U_{eff} parameter, which represents the difference between the Coulomb (U) and exchange (J) parameters $(U_{eff} = U - J)$ is utilised in the DFT+U calculations, in order to model the electron correlation seen in transition metals. (37) A Ueff value of 7.0 eV was applied to Fe3+, determined after comparing experimental structure parameters and properties(44, 45, 46, 47, 48) to those calculated at increasing values of Ueff. (34) A 2x2x1 supercell with antiferromagnetic G ordering of

the Fe³⁺ ions was used for all calculations. The energies of supercells containing dopants and defects were calculated at the lattice parameters of the relaxed perfect supercell.

For studying A- and B-site transition metal dopants, a $U_{\rm eff}$ value of 4.0 eV was used for both Mn^{2+} and Co^{2+} and a value of 7.0 eV was used for Ni^{2+} and Cu^{2+} , these values were evaluated using the method used in ref. [34] to evaluate the $U_{\rm eff}$ value for Fe^{3+} ; comparing experimental parameters of the metal monoxide with those calculated for a range of values of $U_{\rm eff}$, (see Supporting Information) while also taking into account common values used in the literature; between 4 and 5 eV for $CoO_{\circ}^{(44, 49, 50)}$ between 5 and 8 eV for $NiO_{\circ}^{(51-54)}$ 7 eV for $CuO^{(55-58)}$ and between 3 and 5 eV for $MnO_{\circ}^{(53,59-60)}$ when our results showed little structural change between the monoxides at different values of $U_{\rm eff}$.

2.1. Calculating Solution Energies using DFT+U: Solution energies of dopants in charge state q ($\Delta H_f(q)$) were calculated using the following

$$\begin{array}{l} \Delta H_f(q) = E^{dopant} - E^{perfect} \pm \sum n_x (E_x + \mu_x) + \\ q(E_{VBM} + E_F + \Delta E_{pot}) + q^2 E_{ic} \quad , \ (1) \end{array}$$

with E^{dopant} being the total energy of the relaxed supercell, at constant volume, containing the dopant, $E^{perfect}$ the energy of the relaxed perfect supercell, n_x is the number of atoms or ions added or remove from the system, E_x is the elemental reference energy, the energy of an element in its standard state e.g. in a pure metal, and μ_x is the chemical potential of the dopant. When $q \neq 0$, the chemical potential of the electrons in the system needs to be taken into account to correct for the imbalance of charge, which is done with E_{VBM} , the energy of the valance band maximum (VBM), and E_F , the Fermi energy defined with respect to the VBM. In addition, ΔE_{pot} accounts for the difference between the potential of the perfect supercell and the supercell containing the charged dopant, and E_{ic} , a correction to the interaction of charged defects with their periodic image. (61) In this work, E_{ic} has been calculated using the method outlined by Murphy and Hine, (62) as LaFeO₃ is orthorhombic and this method takes into account the differences in dielectric tensors along the a, b and c lattice parameters. For E^{dopant} , one dopant in a 2x2x1 supercell, containing 80 atoms, was used, leading to a doped stoichiometry of La_{0.94}M_{0.06}FeO₃ for A-site dopants and LaFe_{0.94}M_{0.06}O₃ for B-site dopants.

The chemical potential, μ_x , was varied by changing the elemental reference state from a pure metal to a metal oxide, using the anion poor and anion rich (representing the limits of low and high oxygen partial pressures, p_{O_2}) equations respectively. Under anion poor (i.e. reducing) conditions the replaced ion, A, precipitates out as a metal, whereas under anion rich (i.e. oxidising) conditions the replaced ion forms an oxide with the excess oxygen. Here standard Kröger-Vink notation⁽⁶³⁾ has been used (with nickel as an example dopant on the A-site):

$$2A_A^X + O_O^X + 2Ni(s) \leftrightarrow 2Ni_A' + 2A(s) + V_O^{\bullet \bullet} + \frac{1}{2}O_2$$
(2)

$$2A_A^X + O_O^X + 2NiO \leftrightarrow 2Ni_A' + V_O^{\bullet \bullet} + A_2O_3$$
, (3)

For anion poor, nickel metal is used as the elemental reference state and for anion rich nickel(II)oxide is used. The above equations use oxygen vacancies, $V_0^{\bullet\bullet}$, to compensate for the charge imbalance caused when divalent ions replace the trivalent ions on the A- and B-site of LaFeO₃. Here two divalent dopants are introduced, $2Ni_A'$, and the resulting net charge of -2 is balanced with one oxygen vacancy, $V_0^{\bullet\bullet}$. Another compensation

mechanism is through the formation of holes, h^{\bullet} . This mechanism was also investigated in both anion poor and anion rich conditions, which are represented respectively as follows (again using nickel as a demonstrative example):

$$A_A^X + Ni(s) \leftrightarrow Ni_A' + A(s) + h^{\bullet}$$
 (4)
 $A_A^X + NiO + \frac{1}{4}O_2 \leftrightarrow Ni_A' + h^{\bullet} + \frac{1}{2}A_2O_3$. (5)

At ambient temperatures and pressures anion rich represents the more realistic operating conditions for SOFCs.

2.2. Calculating Binding Energies: The extent to which a dopant ion increases the conductivity of LaFeO₃ will depend on both its solution energy and the extent to which the dopant binds the charge compensating species produced. Significant binding energies can lead to a decrease in the conductivity as the mobile ions and electrons can be trapped by the dopant. The binding energies to oxygen vacancies and holes were calculated using the following respective equations:

$$E_{Bind} = E_{cluster} - (E_{V_O^{\bullet \bullet}} + 2E_{M_A'}) \qquad , \quad (6)$$

$$E_{Bind} = E_{cluster} - (E_{h^{\bullet}} + E_{M_A'}) \qquad , \quad (7)$$

where $E_{\rm cluster}$ is the formation energy of the dopant and vacancy or hole cluster, $E_{V_O^{\bullet\bullet}}$ is the formation energy of the isolated oxygen vacancy and $E_{M_A'}$ is the formation energy of the isolated A- or B-site dopant and E_{h^\bullet} is the defect energy of an isolated hole. A negative value indicates that the cluster is more stable than the isolated defects and therefore the dopant will bind to the charge compensating species. As shown in Figure 1, there are two inequivalent oxygen sites in LaFeO₃, situated in the axial (O1) and equatorial (O2) position in the FeO₆ octahedra; binding energies to oxygen vacancies in both sites have been calculated.

To calculate the solution energy of dopant-vacancy clusters, the most stable configuration of two dopants and an oxygen vacancy in a 2x2x1 LaFeO₃ supercell needs to be identified. This was achieved using the Site Occupancy Disorder (SOD) program⁽⁶⁴⁾ which identifies all the unique configurations of a given composition. In this study, the most stable configuration of the dopant-vacancy clusters was established in two steps: initial screening of all unique configurations was performed using interatomic potentials, then the resulting ten most stable configurations were relaxed using DFT+U and the most favourable configuration was found, which was done first with two dopant ions, without the compensating oxygen vacancy, with the process repeated, to find the most favourable site for the compensating oxygen vacancy, using the doped LaFeO₃ supercell. The interatomic potential calculations were performed using the short range potential parameters derived by Cherry et al. (65) for LaFeO₃ and Lewis and Catlow⁽⁶⁶⁾ for the dopant species with the exception of Cu2+ which was derived by Islam and Winch. (67) These calculations were carried out using the GULP(68) code.

3. Results

We will consider the effects of doping the A-site and the B-site separately, first discussing our results for A-site doping.

3.1. A-site: The solution energies of a range of divalent alkaliearth metal and transition metal dopants on the A-site of La-FeO₃ have been calculated with both hole and oxygen vacancy compensation investigated, using equations 1 to 5, as described above. The results are shown in Table 1 in comparison with the experimental metal-oxygen bond distance in the associated bi-

nary oxide, ⁽⁶⁹⁻⁷⁵⁾ a parameter chosen to take into account the effect of Jahn-Teller distortions on the octahedra for copper, which is not represented by ionic radii.

Table 1. The solution energies of divalent alkali-earth metal and transition metal A-site dopants', in anion poor and anion rich conditions, using either hole (h') or oxygen vacancy (V_0 ") compensation, along with the metal-oxygen bond length in the dopants associated binary oxide; M-O. ^aRef 69 ^bRef 70 ^cRef 71 ^dRef 72 ^cRef 73 ^fRef 74 ^gRef 75.

		Solution Energy / eV			
M	M-O/ \mathring{A}^a	h	h*		o "
		Poor	Rich	Poor	Rich
All	kali Earth Met	tals			
Mg_{La}'	2.109^{a}	4.961	1.739	3.598	1.450
Ca _{La} '	2.406^{b}	3.316	0.587	2.501	0.682
Sr_{La}	2.580°	3.306	0.079	2.494	0.343
Ba _{La} '	2.770^{d}	3.992	0.248	2.952	0.456
Ti	ransition Meta	als			
Cu_{La}	1.951 ^e	9.050	1.886	6.324	1.548
Ni_{La}	2.089^{a}	7.762	2.110	5.465	1.697
Co _{La} '	2.130^{f}	7.162	1.343	5.065	1.186
Fe _{La} '	2.163g	5.896	1.551	4.221	1.325
Mn _{La} '	2.223ª	6.159	1.330	4.397	1.177

For the alkali earth dopants, the solution energies decrease with increasing metal to oxygen (M-O) bond length until barium, for which the solution energy increases. This increase in solution energy is due to divalent barium having a larger ionic radius of 1.42 Å, $^{(76)}$ than that of the trivalent lanthanum ion it has replaced (1.36 Å $^{(76)}$). For the transition metals, the solution energies decrease as the M-O bond length increases. For both sets of dopants, the closer their M-O bond length is to the average La-O bond length (2.596 Å) in this material, the lower is its solution energy. This trend is followed by both compensation mechanisms, as is shown clearly in Figure 2.

For the transition metal dopants we should consider the spin orientation of the dopant. In stoichiometric LaFeO $_3$, the B-site Fe $^{3+}$ ions are antiferromagnetically ordered which may affect the spin direction of dopants in either the A- or B-site. Therefore, the solution energies of transition metal dopants in both spin up and down orientations, with respect to the c lattice parameter, were calculated and compared. On the A-site, there was no significant difference observed between these solution energies, the largest difference is observed for Fe $^{2+}$; 0.02 eV, suggesting spin does not play a strong role on this site.

Clearly incorporating dopants under anion rich conditions is more favourable for both sets of dopants and both compensation mechanisms, compared to anion poor conditions, which is driven by the formation of the stable A_2O_3 product under anion rich conditions. Interestingly, under anion rich conditions, (which are more likely to prevail in many experimental conditions), the difference between the solution energies of the two compensation mechanisms is small indicating that, in doped

systems formed under these conditions, both holes and oxygen vacancies will be present, demonstrating the mixed ionic electronic properties of doped systems. Of the alkali earth dopants, strontium has the lowest solution energy, under both conditions. However, under anion poor conditions, the solution energies of strontium and calcium are almost identical, while, oxygen vacancy compensation is clearly more favourable; therefore, incorporating dopants under these conditions will serve to increase the ionic, but not the electronic conductivity of the material

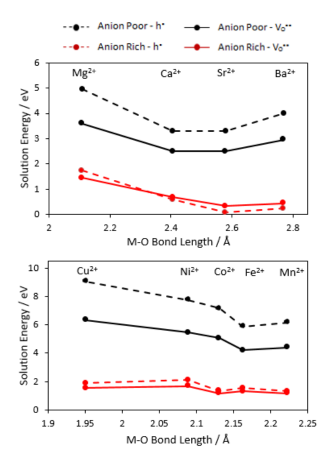


Figure 2. Solution energies of A-site dopants with increasing M-O bond length under anion poor (black) and anion rich (red) conditions; alkali earth metals are on top and transition metals on the bottom, with oxygen vacancy (solid line) and hole (dashed line) compensation.

For the transition metals, manganese has the lowest solution energy, under both anion rich and poor conditions. However, in anion rich conditions, there is only a small difference between the solution energies of all the transition metals studied. As with the alkali earth dopants, there is a significant difference between oxygen vacancy and hole compensation under anion poor conditions, with oxygen vacancies forming in preference, leading to an increase in oxygen vacancy concentration on doping under these conditions, but not holes. Whereas anion rich conditions will promote the formation of both types of compensation.

3.1.1. Hole Localisation: The extent to which a hole localises was determined by comparing the charge densities of the doped and undoped systems. Due to the self-interaction error present in DFT, modelling hole localisation accurately is a challenge

and often produces erroneous results. However, in our investigation we found that introducing a hole only affected the oxygen sub lattice in LaFeO₃ in a minor way; either by the hole localising on a set of specific oxygen ions, or by delocalising throughout the lattice. As only one sub lattice is involved, the most significant error using GGA+U will be systematic, allowing us to study the trends in hole localisation for each set of dopants. (Distorting oxide ions was found not to promote small polaron formation on those ions).

For the alkali earth metal dopants, when a hole is generated in compensation, the hole tends to delocalise around the lattice, with only slight localisation on the oxygen ions surrounding the dopant. This delocalisation increases from magnesium to strontium, before decreasing for barium. For the transition metals, however, the hole is consistently localised on the nearest neighbour oxygen ions surrounding the dopant from manganese to copper. The delocalisation of the hole in the case of alkali earth doping, (except for magnesium) may account for the hole compensation being slightly more favourable than oxygen vacancy compensation under anion rich conditions for these dopants, whereas the opposite is seen for those dopants that show hole localisation.

The effect of hole localisation on the oxygens surrounding the dopant, for Mg and Ba, is further demonstrated by the change in metal-oxygen bond lengths caused by the dopant: the greater the difference from the lanthanum-oxygen bond length, the higher the degree of hole localisation (or small polaron formation). For both sets of A-site dopants, where localisation is observed, the hole predominantly localises on the six oxygens that have the shortest La-O bonds in the pure material. The change in bond lengths on doping for these six bonds are shown in Tables 2 and 3 for the alkali earth and the transition metal dopants respectively, a positive number indicates an increase in bond length, whereas a negative value shows a decrease.

Table 2. The difference between bulk LaFeO $_3$ La-O bond lengths and the M-O bond lengths in alkali earth doped La-FeO $_3$ with hole compensation.

La-O / Å		Difference / Å				
La-O / A	Mg-O	Ca-O	Sr-O	Ba-O		
2.555	1.00	-0.04	0.09	0.23		
2.659	0.83	0.01	0.08	0.16		
2.659	0.83	0.01	0.08	0.16		
2.442	-0.40	-0.01	0.09	0.21		
2.442	-0.40	-0.01	0.09	0.21		
2.407	-0.37	-0.01	0.09	0.19		

For the alkali earth dopants, where hole localisation is only observed in magnesium and barium doped LaFeO $_3$, only these dopants show a notable change in bond length compared to the La-O bond. The localisation of the hole adjacent to Mg^{2+} and Ba^{2+} is due to the significant difference in ionic radii between these dopants and La^{3+} ; the dopants with similar radii; Sr^{2+} and Ca^{2+} , do not cause hole localisation.

For the transition metal dopants, hole localisation on the surrounding oxygens is observed for all dopants, but to varying degrees. The largest hole density on neighbouring oxygen ions is

seen on cobalt and copper, which show the largest difference in bond lengths compared to La-O. Another factor that needs considering when discussing hole localisation for transition metal dopants, is the possibility of the hole localising on the dopant itself, due to their variable oxidation states. The extent of hole localisation on the dopant was established by comparing the relaxed magnetic moment of the dopant, in the 2x2x1 LaFeO₃ supercell, with its expected value as a divalent dopant. The expected values for the transition metal dopants are; 5, 4, 3, 2 and $1 \mu_B$ for Mn, Fe, Co, Ni and Cu respectively. The relaxed magnetic moments have been scaled to account for Fe³⁺, which has a d^5 configuration, having a relaxed magnetic moment of 4.4 μ_B in our supercell. After scaling, the relaxed magnetic moments of the transition metal dopants in an A-site are; 5.1, 4.3, 3.2, 2.1 and $0.85 \mu_B$, in line with those expected for these dopants in their divalent charge state, demonstrating that there is no hole localisation on the transition metal dopants when they occupy the A-site.

Table 3. The difference between bulk LaFeO $_3$ La-O bond lengths and the M-O bond lengths in transition metal doped LaFeO $_3$ with hole compensation.

La-O / Å	Difference / Å					
La-O / A	Mn-O	Fe-O	Co-O	Ni-O	Cu-O	
2.555	0.02	-0.46	1.16	0.62	0.97	
2.659	0.38	0.44	0.94	0.54	0.85	
2.659	0.35	0.44	0.94	0.54	0.73	
2.442	-0.30	-0.43	-0.50	-0.43	-0.42	
2.442	-0.31	-0.43	-0.52	-0.43	-0.52	
2.407	-0.27	-0.45	-0.54	-0.40	-0.42	

3.1.2. Binding Energies: The binding energies of these divalent dopants with the two charge compensating species should be considered before choosing the most appropriate dopant. All dopants studied promote both oxygen vacancy and hole formation. However, to effect an increase in conductivity, the dopant must not bind strongly to these species. Tables 4 and 5 show the binding energies for the divalent A-site alkali earth and transition metal dopants respectively.

Table 4. Calculated binding energies of A-site alkali earth dopants with oxygen vacancies and holes in LaFeO₃, binding energies for oxygen vacancies in both the O1 and O2 sites have been calculated.

Domant	F	Binding E	nergy / eV	V
Dopant	Mg	Ca	Sr	Ba
V _{O1} "	-0.393	-0.034	-0.061	-0.620
V_{O2} "	-0.102	-0.033	0.092	-0.507
h*	0.243	0.202	0.188	0.129

Oxygen vacancies do not bind significantly to either calcium or strontium: the weak binding of calcium, and strontium to O1 vacancies, will be overcome under SOFC operating conditions

(kT = 0.075 eV at 873 K, the lower limit of intermediate temperature SOFC operating temperatures). However, magnesium and particularly barium have larger binding energies, very probably due to the mismatch in ionic radii of these dopants compared to lanthanum (0.89, 1.61 and 1.36 Å respectively for Mg^{2+} , Ba^{2+} and La^{3+}), as has been observed in related systems. (77) It is of interest that all dopants have a stronger binding to vacant O1 sites compared to O2 sites, a promising result as O2 vacancies form more favourably compared to O1 vacancies.

The site of the oxygen vacancy in each doped system, which was determined separately for each dopant using SOD(64), is dependent on the dopant species as well as the type of oxygen vacancy; O1 or O2, causing the difference in the binding energies observed for the two sites. For the O1 site, with the exception of magnesium, the vacancy occupies one of the nearest neighbour oxygen sites to one of the dopants, whereas for the O2 site, for strontium and barium, the vacancy is in an oxygen site between the two dopants, which distributes the charge from the vacant site between the two dopants, causing a smaller binding energy to either of the positively charged dopants. The exceptions to this are magnesium - for which the O2 vacancy is located next to only one dopant and the O1 site is not a nearest neighbour site, but a next nearest neighbour site, and calcium, for which both O1 and O2 sites are bonded only to one dopant, which leads to the similarity observed between the binding energies of calcium to the two oxygen vacancy sites.

None of the alkali earth metal dopants studied bind to holes, suggesting these dopants should cause an increase in electrical conductivity in this material, if incorporated under the appropriate conditions. Combining these results with the solution energies calculated suggests that strontium or calcium would be appropriate A-site alkali earth metal dopants, in line with experimental results which show increased ionic conductivity in Srand Ca- doped LaFeO₃ compared to the undoped material, (19,23) as well as an increase in the electronic conductivity, . (78, 79, 80, 81)

Table 5. Calculated binding energies of A-site transition metal dopants with oxygen vacancies and holes in LaFeO₃, binding energies to oxygen vacancies in both the O1 and O2 sites have been calculated.

		Bindi	ng Energy	y / eV	
Dopant	Cu	Ni	Co	Fe	Mn
V _{O1} "	-0.380	-0.567	-0.054	-0.464	-0.143
V_{O2}	-0.159	-0.624	-0.112	-0.117	-0.155
h*	0.303	0.284	0.237	0.154	0.224

Interestingly, whereas the alkali earth metal dopants bind to O1 oxygen vacancies stronger than to O2, the transition metal dopants have a stronger binding to O2 oxygen vacancies, in the majority of cases, the key exceptions being Fe and Cu. The binding energy of transition metal dopants to oxygen vacancies generally increase as the M-O bond length of the associated binary oxide decreases, except for Cu, which shows a lower binding energy than expected and Fe with an O1 vacancy, which is significantly higher than expected. The difference for copper can be attributed to the greater stability of Cu in a square planar geometry compared to the 12-coordinate geometry of lanthanum in LaFeO3. The A-site copper dopant shifts away from the La site into a square planar site between four of the oxygen ions,

stabilising the dopant and decreasing the magnitude of its binding energy, which can be seen in Figure 3. However, this distortion to the lattice makes it highly unlikely that copper will occupy an A-site in this material, and favours the B-site instead, a point discussed in the next section.

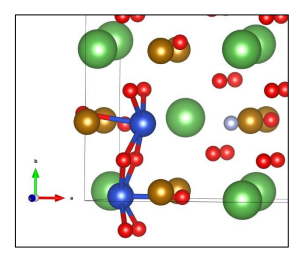


Figure 3. The most stable dopant-vacancy cluster for an A-site copper dopant, showing relaxed position of an A-site copper dopant (blue). The dopant has shifted significantly away from one oxygen ion to form a pseudo-square planar geometry with four of the other oxygen ions. This shift will be more favourable if the oxygen site it is shifting away from is vacant, as is shown here with the oxygen vacancy shown in grey.

The significantly larger binding energy for Fe to an O1 oxygen can be explained as due to the nucleation of a secondary Fe_2O_3 phase within the structure, seen in Figure 4. As the O2 vacancy is a nearest neighbour to one of the Fe ions it breaks up this secondary phase; therefore, this higher binding energy is only seen for the O1 vacancy. The tendency of the majority of transition metal dopants to bind to O2 site vacancies is probably due to the closer proximity of the O2 site to the lanthanum site compared to O1.

As with the alkali earth dopants, none of the transition metal dopants studied bind significantly to holes. Combining these results with the calculated solution energies suggests that Mn^{2+} could be an appropriate A-site dopant, due to its low solution energy and small binding energy.

Although strontium is often used as an A-site dopant for perovskite materials such as LaFeO₃, ^(5, 82, 83) manganese has previously been assumed to occupy the B-site when used as a dopant, a point that is discussed further in the next section. It would be of interest to investigate the effect of doping the A-site with both strontium and manganese which, according to our results, would lead to a promotion in both of bulk ionic and electronic conductivity in LaFeO₃.

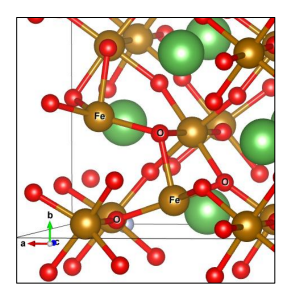


Figure 4. The high binding energy of Fe to O1 vacancies indicates incipient nucleation of an Fe_2O_3 phase. The A-site Fe dopants are labelled Fe and the oxygens labelled O are oxygens with a coordination number of 3, a feature of the Fe_2O_3 lattice. The O1 vacancy is shown in grey.

3.1.3. Comparison of Site Stability: The promising results displayed by manganese as an A-site dopant leads to the need to establish the likelihood of manganese occupying an A-site compared to a B-site in LaFeO₃. The Goldschmidt tolerance factor, t, (84) is a hard sphere model that uses ionic radii to assess the distortion of the lattice caused by a dopant ion. The larger the distortion the less likely the dopant is to occupy that lattice site. It is calculated as follows

$$t = \frac{r_x + r_A}{\sqrt{2} (r_x + r_B)}$$
 , (8)

where r is the ionic radii of the ion with A being the A-site cation, B being the B-site cation and X being the anion; oxygen in this case. An ideal cubic perovskite would have a Gold-schmidt factor of 1 with the lattice becoming more distorted the more the Goldschmidt factor deviates from 1. The Goldschmidt tolerance factor for pure stoichiometric LaFeO₃ is 0.908, for comparison. To calculate the Goldschmidt factor of the doped system, the ionic radius of the divalent dopant was used instead of the ionic radii of La³⁺ or Fe³⁺ for A-site and B-site dopants respectively; these are shown in Table 6.

The radius of an ion is highly dependent on the ion's coordination number in the material, with the ionic radius increasing with the coordination. For the transition metals only ionic radii up to a coordination number of 8 are available, so that although the tolerance factor values calculated for the B-site dopants will be accurate – using a coordination number of 6 – those calculated for the A-site will be less so, as the ionic radii associated with a coordination of 8, or 6 for nickel and copper, rather than the actual value of 12, were used. However, for the A-site, the tolerance factor increases with larger ionic radii. As the ionic radius is expected to increase with a higher coordination number, we expect a higher tolerance factor, and therefore the predicted stability of the perovskite structure will be greater.

It is clear that the majority of the transition metal dopants favour the B-site, causing less distortion compared to when they occupy the A-site, even with the underestimation of the A-site tolerance factors. However, manganese only causes slightly less

distortion on the B-site compared to the A-site, especially considering this value could be underestimated. This similarity suggests that it may be possible to form LaFeO₃ with manganese as an A-site dopant, an interesting avenue of further investigation.

Table 6. The Goldschmit tolerance factor of divalent dopants on the A-site and B-site of LaFeO₃.

Donant	Goldschmidt Factor			
Dopant	A-site	B-site		
Alkali Eartl	n Metals			
${ m Mg_{Fe}}'$	0.734	0.876		
Transition	Metals			
Cu _{Fe} '	0.737	0.872		
Ni _{Fe} '	0.722	0.889		
Co _{Fe} '	0.797	0.865		
Fe _{Fe} '	0.804	0.851		
$Mn_{Fe}{^{\prime}}$	0.818	0.832		

3.2. B-site: The solution energies of a range of divalent transition metal dopants on the B-site of LaFeO₃ have been calculated with both hole and oxygen vacancy compensation. Magnesium, as the smallest alkali earth metal, was also considered as a dopant on the B-site for comparison.

Considering the transition metal dopants initially, we see that, as with the A-site dopants, anion rich provides the most favourable formation conditions for both compensation mechanisms. However, unlike the A-site dopants, the orientation of the spin on the transition metal dopant ion was found to have an impact on the solution energy of the dopants, with lower solution energies if the spin orientation of the dopant is in line with the antiferromagnetic ordering of the Fe^{3+} ions – a dopant replacing a spin down Fe^{3+} would favourably be in a spin down orientation.

Table 7. The solution energies of divalent B-site metal dopants in anion poor and anion rich conditions, using either hole (h') or oxygen vacancy (V_0 ") compensation along with the metal-oxygen bond length in the dopants associated binary oxide; M-O. ^aRef 69 ^bRef 73 ^cRef 74

			Solution E	inergy / eV	T .
M	M-O/ Å	h*		V	o "
		Poor	Rich	Poor	Rich
Alkali Earth Metals					
Mg_{Fe}'	2.109 ^a	0.737	1.038	0.784	0.985
Tr	ansition Met	als			
Cu_{Fe}	1.951 ^b	4.872	1.283	3.538	1.146
Ni_{Fe}	2.089^{a}	2.938	0.861	2.249	0.865
CoFe'	2.130^{c}	2.891	0.648	2.218	0.722
$Mn_{Fe}{^{\prime}}$	2.223a	2.307	1.053	1.828	0.993

Interestingly, different trends are observed for the two conditions investigated. Under anion poor conditions the solution en-

ergies decrease with increasing M-O bond length. Whereas, under anion rich conditions the solution energies reach a minimum at cobalt and then increase again. As with transition metal dopants on the A-site, there is only a small variation in the solution energies of the dopants studied under anion rich conditions, with the solution energies of dopants with the different compensation mechanisms being comparable so that both holes and oxygen vacancies are likely to be present in B-site doped LaFeO₃ formed under this condition. Under anion poor, however, oxygen vacancies are more favourable.

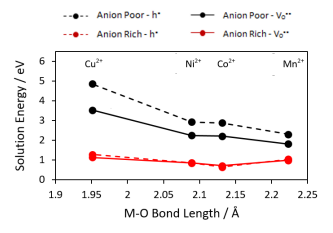


Figure 5. Solution energies of B-site transition metal dopants with increasing M-O bond length under anion poor (black) and anion rich (red) conditions. Solution energies are calculated with oxygen vacancy (solid line) and hole compensation (dashed line).

3.2.1. Hole Localisation: As with the A-site dopants, only the oxygen sub lattice is affected when a hole is introduced in compensation, and the self-interaction error in DFT largely cancels. As a consequence, we can analyse the trends in hole localisation observed for these dopants.

Hole localisation with the B-site transition metal dopants follows a similar trend to the A-site alkali earths. Initially, for manganese, the hole is localised on the six surrounding oxygen ions; this localisation decreases from manganese to cobalt, before increasing again for nickel and copper.

The oxidation state of each of the dopants was established via their magnetic moment, following the same method used for the A-site transition metal dopants. It was found that all dopants are in their 2+ charge state, having magnetic moments of; 4.7, 3.0, 1.95 and 0.83 μ_B respectively for Mn, Co, Ni and Cu. Therefore, no localisation is expected on the dopant.

As with the A-site dopants, the hole localisation can be identified by the effect the dopant has on the bond length between the B-site ion and the oxygen ions. The greater the change in bond length caused by the dopant, shown in Table 8, the more localised the hole is on the surrounding oxygens, as the change in bond length is caused by the localised polaron. The largest difference in bond length is seen when manganese is the dopant, and oxygen ions surrounding manganese show the highest degree of localisation. The change in bond length then decreases for cobalt, with almost no change in bond length observed, consistent with no localisation seen on the oxygen ions surrounding

cobalt. Finally, the change in bond length increases again for Ni and Cu

Table 8. The difference between bulk LaFeO₃ Fe-O bond lengths and the M-O bond lengths in transition metal doped LaFeO₃ with hole compensation. Those shown in bold are the bond lengths with oxygen ions that show hole localisation.

Fe-O / Å		Differe	nce / Å	
Fe-O / A	Mn-O	Co-O	Ni-O	Cu-O
2.023	0.01	0.00	0.03	0.06
2.028	0.09	0.00	0.04	0.03
2.018	0.05	0.01	0.04	-0.02

3.2.2. Binding Energies: The binding energies for the divalent B-site dopants to oxygen vacancies and holes are shown in Table 9.

The results show that all B-site dopants studied bind to both oxygen vacancies and holes. These dopants may therefore decrease the ionic and electronic conductivity of LaFeO₃; however the incorporation of B-site dopants could be necessary for the stability of the material and therefore a dopant should be chosen that limits this detrimental effect. Of the transition metals studied, nickel has the lowest binding energy for both oxygen vacancies and holes, although magnesium has lower binding energies to both compensating species. This observation would lead to the conclusion that cobalt, nickel and magnesium would be the good B-site dopants, as they have low solution energies and the minimal binding energies to both oxygen vacancies and holes promoting ionic and electronic conductivity.

Table 9. Calculated binding energies of B-site dopants with oxygen vacancies and holes in LaFeO₃.

M		Bindi	ng Energy	y / eV	
IVI	Mg	Cu	Ni	Co	Mn
Voi"	-0.16	-0.77	-0.19	-0.35	-0.42
V_{O2} "	-0.17	-1.40	-0.20	-0.42	-0.40
h*	-0.03	-0.25	-0.05	-0.36	-0.68

Experimentally, promising results have been observed when using cobalt and nickel as B-site dopants. LSCF based materials, as mentioned previously, have high ionic and electronic conductivities, as well as being more resistant to chromium poisoning than other cathode materials. $^{(4,5,7,8)}$ La_xSr_{1-x}Fe_yNi_{1-y}O_{3-\delta} (LSNF) on the other hand, has attracted attention as a cathode material for solid oxide electrolysis cells, the reverse of a solid oxide fuel cells, due to its high electronic conductivity, thermal expansion coefficient similar to that of YSZ, and its high coking tolerance to CO. $^{(85,86)}$ These first two properties have resulted in Ni-based perovskites also being considered as cathode materials for SOFCs, $^{(87)}$ and our results indicate that they should have ionic and electronic conductivities comparable to LSCF.

These observed properties suggest that the binding energies calculated in this work are not large enough to inhibit conductivity under operating conditions, particularly when A-site dopants are present. There is also evidence that transition metal

dopants increase the ionic conductivity without A-site dopants, for example Kharton et al.⁽⁸⁸⁾ found doping LaFeO₃ with nickel increased the electrical conductivity and bulk ionic conductivity of the material, demonstrating that the binding energies can be overcome.

The results for magnesium as a B-site dopant show promise: it has a lower solution energy than the transition metal dopants and binds less to both oxygen vacancies and holes. In addition, the tolerance factors for magnesium, shown in Table 6, show that it will cause less strain on the structure of LaFeO3 if situated on the B-site as opposed to the A-site. Therefore magnesium, with its low solution energy and low binding energy to both oxygen vacancies and holes, could be a promising B-site dopant. This has recently been demonstrated experimentally; when comparing LaFeO3 to LaFe1-xMgxO3 (x = 0.05) Díez-García and Gómez (89) found that the doped species had a higher efficiency as a photocathode for the oxygen reduction reaction, a key reaction in SOFCs.

The current work has only focused on solution energies and binding energies of these dopants. Other factors will play a role in the suitability of these dopants, including the effect they have on the thermal expansion coefficient (TEC). The effect of nickel and cobalt on the TEC of LaFeO₃ is well reported; the presence of cobalt as a dopant causes as increase in the TEC, usually linearly with cobalt content, (90) so low concentrations of cobalt are often used in order to keep the TEC similar to those of common electrolyte material, whereas Ni-doped LaFeO₃ materials have favourable TECs, in line with those of common electrolyte materials, even at large concentrations. (91,92) The effect of Mg on the TEC of LaFeO3 is less well reported, and would need to be investigated before proposing Mg as a B-site dopant. However, results from Mg B-site doping of a related compound, La_{0.7}Sr_{0.3}MnO_{3-δ} are promising, with low TECs and improved electrochemical performance compared to LaSrMnO3 reported.(93)

Comparing the solution energies of manganese on both the A-site and the B-site; under anion poor conditions the solution energies of manganese as a B-site dopant are significantly smaller compared to its solution energy as an A site dopant. However, under anion rich conditions, the solution energies are very similar, particularly for oxygen vacancy compensation for which the solution energy of manganese on the B-site is only 0.045 eV lower than that of the A-site.

3.2.3 Variable Oxidation States: Transition metals are able to occupy a variety of oxidation states within a material, a factor which can help or hinder the conductivity properties. With this in mind, we calculated the barrier to oxidation of the appropriate divalent transition metal B-site dopants in LaFeO₃. Manganese, cobalt, nickel and copper could form 3+ (or 4+ for manganese) oxidation states within LaFeO₃, therefore oxidation to these states was considered.

The barrier to oxidation was calculated based on the following equation, using manganese as an example dopant;

$$Fe_{Fe}^X + Mn_{Fe}' \leftrightarrow Mn_{Fe}^X + Fe_{Fe}'$$
 (9)

With an addition oxidation step considered for generating manganese 4+

$$Fe_{Fe}^X + Mn_{Fe}^X \leftrightarrow Mn_{Fe}^{\bullet} + Fe_{Fe}^{\prime}$$
 (10)

The reaction energies are shown in Table 10.

The calculated energies show that, particularly for manganese, there is a low barrier to oxidising these divalent dopants

which are therefore likely to be present in 2+, 3+ and 4+, for manganese, oxidation states, in agreement with experimental findings. (94) However, nickel, not surprisingly, has a high barrier to oxidation. Interestingly, whereas the other dopants favour high spin configurations, nickel favourably oxidises into the low spin state.

Table 10. The solution energies of transition metal B-site dopants' undergoing oxidation to form 3+ oxidation states and 4+ in the case of manganese. In all cases the dopants in both 2+, 3+ and 4+ states are in high spin states, the value for nickel in brackets represents the oxidation of nickel into a low spin state.

M	Solution Energy / eV
Mn_{Fe}^{X}	0.300
Mn_{Fe}	0.735
Co_{Fe}^X	0.860
Ni_{Fe}^{X}	2.190 (1.782)
$Cu_{Fe}{}^{X} \\$	1.145

4. Summary and Conclusion

The most appropriate A- and B-site dopants were evaluated from a range of divalent transition metals with divalent alkali earth metals also investigated for the A-site. We find for A-site alkali earth metal doping that there is only minimal binding to oxygen vacancies and holes, with the exception of barium which binds strongly to oxygen vacancies in both sites. Low solution energies were calculated for Ca²⁺, Sr²⁺ and Ba²⁺ under anion rich conditions, with Sr²⁺ and Ca²⁺ emerging as the most appropriate due to a combination of low solution energy and minimal binding energies.

For A-site transition metal dopants, the solution energies are all comparable. However, manganese has low binding energies compared to other transition metal A-site dopants, which combined with the minimum distortion of the surrounding LaFeO $_3$ lattice suggests that it would be the most appropriate transition metal A-site dopant in LaFeO $_3$. Comparison with manganese as a B-site dopant showed comparable solution energies under oxygen rich conditions and similar Goldschmidt tolerance factors, suggesting manganese could be a viable A-site dopant, in contrast to its widely assumed B-site position. Cobalt nickel and magnesium were found to be the most favourable B-site dopants, due to low solution energies and low binding energies to oxygen vacancies, along with minimal distortion - particularly for nickel — to the LaFeO $_3$ lattice.

Under anion rich conditions, all dopants, in both sites, demonstrate similar solution energies between the two charge compensation mechanisms investigated; oxygen, vacancies and holes. This result is particularly significant as it explains the origins of the mixed ionic electronic conductivity observed in doped perovskites, such as LSCF, a property that is vital to the application of these materials as cathode materials in intermediate temperature SOFCs.

ASSOCIATED CONTENT

Supporting Information. U-Values for Metal Oxides.

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Author Contributions

The manuscript was written through contributions of all authors.

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