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# Factors Governing Oxygen Vacancy Formation in Oxide Perovskites

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**ABSTRACT:** The control of oxygen vacancy ( $V_0$ ) formation is critical to advancing multiple metal-oxide-perovskite-based technologies. We report the construction of a compact linear model for the neutral  $V_0$  formation energy in ABO<sub>3</sub> perovskites that reproduces, with reasonable fidelity, Hubbard-*U*-corrected density functional theory calculations based on the state-of-the-art, strongly constrained and appropriately normed exchange-correlation functional. We obtain a mean absolute error of 0.45 eV for perovskites stable at 298 K, an accuracy that holds across a large, electronically diverse set of ABO<sub>3</sub> perovskites. Our model considers perovskites containing alkaline-earth metals (Ca, Sr, and Ba) and lanthanides (La and Ce) on the A-site and 3*d* transition metals (Ti, V, Cr, Mn, Fe, Co, and Ni) on the B-site in six different crystal systems (cubic, tetragonal, orthorhombic, hexagonal, rhombohedral, and





monoclinic) common to perovskites. Physically intuitive metrics easily extracted from existing experimental thermochemical data or via inexpensive quantum mechanical calculations, including crystal bond dissociation energies and (solid phase) reduction potentials, are key components of the model. Beyond validation of the model against known experimental trends in materials used in solid oxide fuel cells, the model yields new candidate perovskites not contained in our training data set, such as  $(Bi,Y)(Fe,Co)O_3$ , which we predict may have favorable thermochemical water-splitting properties. The confluence of sufficient accuracy, efficiency, and interpretability afforded by our model not only facilitates high-throughput computational screening for any application that requires the precise control of  $V_O$  concentrations but also provides a clear picture of the dominant physics governing  $V_O$  formation in metal-oxide perovskites.

#### INTRODUCTION

Metal-oxide perovskites have many applications in fields ranging from energy science to electronics: e.g., two-step thermochemical cycles for water and carbon dioxide splitting (WS and CDS, respectively),<sup>1-7</sup> thermochemical energy storage, <sup>8–11</sup> thermochemical  $O_2$  storage/pumping,<sup>12–14</sup> thermochemical air separation,<sup>14–18</sup> clean electricity production using solid oxide fuel cells (SOFCs),<sup>19-23</sup> solid oxide electrolyzers that produce hydrogen and carbon monoxide from high-temperature electrochemical WS and CDS,<sup>24–26</sup> and ferroelectric random-access memory (FRAM).<sup>27-29</sup> These applications depend sensitively on the concentration of oxygen vacancies (V<sub>O</sub>s), which are capable of dictating thermodynamic, electronic, and emergent properties. As examples, Vos reduce  $H_2O$  to  $H_2$  in a two-step thermochemical WS (often referred to as solar thermochemical hydrogen or STCH),<sup>2,30,31</sup> absorb O<sub>2</sub> to maintain low O<sub>2</sub> partial pressures in thermochemical O<sub>2</sub> storage/pumping and air separation,  $^{12-14}$  facilitate O<sup>2-</sup> transport in SOFCs,  $^{22,23,32,33}$  and pin ferroelectric domain walls that impede ferroelectric switching in FRAM.<sup>34–37</sup> The concentration of V<sub>o</sub>s in any given oxide chiefly depends on their formation energy  $(E_v)$ , which is a wellvetted microscopic descriptor of performance, especially in applications such as STCH<sup>38-40</sup> and SOFCs.<sup>22,23,33,41</sup> The ease with which a neutral  $V_O$  forms in metal (M) oxides, where a neutral  $V_O$  refers to the removal of an entire oxygen atom (in its neutral, ground-state  ${}^{3}P_{2}$  electron configuration) that contributes to half of an O<sub>2</sub> gas molecule (in its neutral, ground-state  ${}^{3}\sum_{g}$ ), leaving behind two electrons from the O<sup>2-</sup> that are donated back to the lattice (typically the cation sublattice), can be described by the following simple physical picture, similar to an extended Born–Haber cycle (see Figure 1).<sup>22,23</sup> First, forming the V<sub>O</sub> breaks bonds (*O*–*M* bond dissociation). Second, the neutral V<sub>O</sub> may reduce its neighboring cations (*M* reduction) or, in materials with aliovalent substitution (e.g., Sr-substituted LaMO<sub>3</sub>), may fill partial holes in the O sp band (*lattice reduction*).<sup>42–47</sup> Third, intrinsic stability/instability can predispose the metal oxide to form fewer/more V<sub>O</sub>s (metal oxide stability).<sup>48–50</sup> Note that stability typically is assessed as the energy above the convex hull (*E*<sub>hull</sub>), which is the energy of

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**Figure 1.** Picturing neutral  $V_O$  formation in metal (M) oxides as an extended Born–Haber thermodynamic cycle. With the metal oxide in its ground-state (gs) polymorph as the starting point, neutral  $V_O$  formation involves O–M bond dissociation typically followed by M reduction. Metastability (ms) of the underlying metal oxide can lead to a decrease in the neutral  $V_O$  formation energy.

decomposition of a given material into the set of most stable materials at its chemical composition.

To better understand the nuances of and optimize  $V_O$  formation in metal oxides, we need deeper understanding of crystal and electronic structure–property relationships. To this end, many have turned to density functional theory (DFT) based calculations coupled with phenomenological model building.<sup>33,51-69</sup> The literature contains a number of models capable of predicting  $V_O$  formation energies: e.g., those based on the O 2*p* band center descriptor.<sup>33,51,62-66</sup> It is important, however, to contrast the strengths of these models with their limitations, in order to illustrate our motivation for building an improved model in terms of physical intuition and ease of calculation.

In terms of strengths, these models generally are quite accurate. For example, Lee et al. introduced the O 2p band center as a descriptor for the Vo formation energy and applied it to the discovery of new multinary perovskites for SOFCs that were verified experimentally.<sup>33</sup> Deml et al. reported a mean absolute error (MAE) of ~0.2 eV for 45 insulating binary and ternary metal oxides using computable quantities such as formation enthalpies, electronic band centers, and Pauling electronegativities.<sup>66</sup> Pavone et al.<sup>23</sup> showed, and Maiti et al.<sup>5</sup> later confirmed, that atomic properties, such as ionization energies and bond dissociation enthalpies, can be used to make predictions in perovskite crystals with surprising accuracy, given the dissimilarity between the chemical environments of atoms/molecules and solid crystals. Most of these models are simple, i.e., they contain only a few terms or descriptors, which suggests that V<sub>O</sub> formation energies are readily describable. While some of the prior model predictions have been validated experimentally, leading to the discovery of new materials for a variety of applications,<sup>33,52</sup> significant progress is still required in the case of applications such as STCH (driven directly by solar or potentially other sources of renewable heat) to achieve improved efficiencies.

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Despite their strengths, the reported models are still limited in a number of ways. The first limitation has to do with the level(s) of DFT used for parametrization. A few models<sup>52,59</sup> have been constructed using the generalized gradient approximation (GGA) exchange-correlation (XC) functional of Perdew, Burke, and Ernzerhof (PBE).<sup>70</sup> While PBE XC can describe alkaline-earth and La oxides fairly accurately (vide infra), this method fails for 3d-transition-metal oxides, where a more accurate treatment of XC is essential.<sup>71</sup> The most widely used method is PBE+U, which pairs the PBE XC functional with Hubbard U corrections to reduce the large self-interaction error incurred by highly localized d electrons in transitionmetal cations.<sup>71</sup> However, the state-of-the-art XC functional for solids is the strongly constrained and appropriately normed (SCAN) meta-GGA XC functional.<sup>72,73</sup> We recently showed that an accurate reproduction of the properties of transitionmetal oxides using the SCAN functional still requires Hubbard U corrections:  $^{74-76}$  i.e., a SCAN+U approach. Our methodology, in which we fit U to relevant oxidation energies, 77,78provides superior predictions of bulk thermodynamics, band gaps, and magnetic structures in comparison to PBE, PBE+U, and SCAN, thus highlighting the usefulness of a database of SCAN+ $UV_{0}$  formation energies that we present herein.

The second major limitation of the existing models is that they, to the best of our knowledge, were trained on homogeneous subsets of perovskite chemical space. Specifically, they lack a diverse sampling of A- and/or B-site metal cations (# unique A × # unique B  $\leq$  10), <sup>33,57,59–61,64,65</sup> space groups (# unique = 1), <sup>33,51,52,59,60,63–65</sup> and electronic structures (i.e., metals, semimetals, and nonmetals).<sup>66,69</sup> The only exception is the study by Maiti et al., <sup>52</sup> which uses the less accurate PBE XC functional for 3*d*-transition-metal-containing perovskite oxides. The lack of diversity in the training data can significantly limit the predictive power of such models, especially when they are extrapolated to higher-component (e.g., quaternary, quinary, etc.<sup>7</sup>) and structurally diverse (e.g., experimentally accessible metastable polymorphs<sup>79–81</sup>) metaloxide chemical spaces.

Finally, model-building approaches that use the band center or band gap as descriptors often rely on computationally expensive hybrid functional or GW calculations to obtain accurate predictions, which makes it hard to train a large data set. Moreover, band-center-based descriptors lack the physical intuition with respect to the physical process of  $V_0$  formation (i.e., M–O bond dissociation, reduction, and stability), as the band *edges* normally play a more significant role than the band *center*. Thus, the need still exists to build a computationally inexpensive yet sufficiently accurate, more physically intuitive, and broadly trained predictive model, especially for the discovery of higher-component or structurally diverse materials.

In this work, we utilize the state-of-the-art SCAN+U method to construct a large database of V<sub>o</sub> formation energies in ABO<sub>3</sub> perovskite oxides for a diverse collection of metal cations, i.e., A-sites (Ca, Sr, Ba, La, and Ce) and B-sites (Ti, V, Cr, Mn, Fe, Co, and Ni), and space groups (cubic, tetragonal, orthorhombic, hexagonal, rhombohedral, and monoclinic), as well as a range of electronic structures, from wide-band-gap insulators and semiconductors to metallic systems. As such, we have set out to provide an accurate (i.e., using SCAN+U), extensive (341 unique V<sub>o</sub>s), and diverse (233 composition–structure combinations) catalog of V<sub>o</sub> formation energies. Subsequently, we fit a theoretical model, using machine-learning techniques



**Figure 2.** ABO<sub>3</sub> perovskite structures. Elements (panel a, non-gray rectangles) and crystal systems (panel b) considered in this study. A-sites (blue and orange) and B-sites (green) are as follows: Ca (2+), Sr (2+), Ba (2+), La (3+), Ce (3+ or 4+), Ti (3+ or 4+), V (3+ or 4+), Cr (3+ or 4+), Mn (3+ or 4+), Fe (3+ or 4+), Co (2+, 3+, or 4+), and Ni (2+, 3+, or 4+).

with simple and physically intuitive descriptors, such as crystal bond dissociation energies and crystal (i.e., solid phase) reduction potentials, to understand and predict  $V_O$  formation energies in a diverse set of metal-oxide perovskite frameworks.

The structure of this paper is as follows. First, we detail the computational methods used. Second, we explain the construction of our model and report its performance, key metrics, and trends for various subsets of the data. Third, we introduce our descriptors for  $V_O$  formation energies and explain how they improve upon the palette of descriptors used in the literature. Fourth, we outline avenues for future research, both in terms of improving our methodology and for applying it to more complex systems (e.g., multinary metal-oxide perovskites and nonoxides). We also discuss the implications of our model with a focus on materials design and candidate identification for various applications. Finally, we conclude by recapitulating our findings and their implications for the understanding and control of  $V_O$  formation in metal-oxide perovskites.

#### COMPUTATIONAL METHODS

Quantum Mechanics Calculations. We performed DFT+ $U^{74-76}$  calculations of neutral V<sub>0</sub> formation energies, crystal reduction potentials, phase diagrams, and band gaps using the Vienna Ab initio Simulation Package (VASP version 5.4.4).<sup>82,83</sup> We used the SCAN XC density functional,<sup>72</sup> with optimal U values taken from previous work.<sup>77,78</sup> To apply the Hubbard U correction, we employed the rotationally invariant scheme of Dudarev et al.<sup>84</sup> as implemented in VASP. We find that SCAN+U calculations are only modestly more computationally expensive (~1.85-2.35×) for each self-consistentfield loop in comparison to those with PBE+U (see Table S1 in the Supporting Information). We utilized the all-electron, frozen-core, projector augmented-wave (PAW) potentials<sup>85,86</sup> as in our prior work<sup>7,77,78,87</sup> to describe the ion-electron interactions, along with including the nonspherical contributions related to the gradient of the electron density and the kinetic energy density within the PAW spheres for the XC evaluation. We used the Accurate "precision" mode in VASP to avoid aliasing errors when the fast-Fourier-transform and support grids were set, employed an additional support grid for more accurate evaluation of augmentation charges, and evaluated the

projection operators in real space using VASP's fully automatic optimization scheme. We used a plane-wave kinetic energy cutoff of 520 eV,  $^{7,77,78,87}$  as this threshold converges total energies of oxide perovskites to within 1 meV/atom.

To perform Brillouin-zone integrations, we used a  $\Gamma$ -point-centered Monkhorst-Pack<sup>88</sup> k-mesh with spacing between k-points of  $\leq$ 0.025 Å<sup>-1</sup>, as previously optimized,<sup>77</sup> and added a Gaussian smearing function with a width of 0.05 eV. For electronic minimizations, we used the blocked Davidson iteration scheme or the conjugate gradient algorithm,<sup>89</sup> depending on difficulties in convergence. In all cases, we deemed to have reached self-consistentfield convergence when the energy change between steps was  $\leq 1 \times$ 10<sup>-5</sup> eV/cell. We performed collinear, spin-polarized calculations and initialized the atomic magnetic moments in a ferromagnetic configuration.<sup>39,90</sup> We considered the high-spin states for Ti, V, Cr, Mn, Fe, Co, and Ni, while for Co and Ni, we also considered the intermediate- and low-spin states. We used the conjugate gradient algorithm to perform structure optimization, stopping the process when the norms of all the atomic forces reached  $\leq |0.03| \text{ eV/Å}$ . For primitive cells, we optimized the atomic positions, cell shape, and cell volume, whereas for supercells, we only optimized the atomic positions to simulate the formation of V<sub>O</sub>s in the dilute limit, in which the bulk lattice structure is not expected to change. Although we preserved symmetry in primitive cell structure optimizations to ensure that there were no symmetry-breaking relaxations (e.g., cubic  $\rightarrow$ tetragonal), we did not preserve the symmetry of ionic positions in supercell calculations, in order to allow for local symmetry breaking upon Vo formation.

**ABO<sub>3</sub> Perovskite Structures.** Figure 2 displays the set of elements (Figure 2a) and structures (Figure 2b) that we considered in constructing our V<sub>O</sub> formation energy data set. On the A-site, we considered alkaline-earth metals (Ca, Sr, and Ba) and lanthanides (La and Ce) that are reasonably abundant and are most likely to adopt the perovskite crystal structure (as opposed to ilmenite-like phases).<sup>91</sup> Of the lanthanides, we focused on La and Ce because they comprise the La<sub>1-x</sub>SrMO<sub>3</sub> (M = Cr, Mn, Fe, Co) family of perovskite-type materials for SOFC applications and CeO<sub>2</sub>, the state-of-the-art metal oxide for STCH, respectively. On the B-site, we considered the redoxactive 3d transition metals (i.e., Ti, V, Cr, Mn, Fe, Co, and Ni), with the exception of Cu, since Cu oxides typically will not survive nominal STCH temperatures. SCAN/SCAN+U also inaccurately describes Cu redox chemistry.<sup>78,92</sup> We did not consider the 4d and 5d transition metals for two reasons, both of which we hope to address in future



Figure 3. SCAN+*U*-calculated vs model-predicted  $E_v$  for (a) room-temperature-stable ( $E_{hull} \le 298.15k_B$ ) and (b) all ABO<sub>3</sub> perovskite structures considered.  $\sum E_b$  is the crystal bond dissociation energy sum in eV,  $V_r$  is the maximum crystal reduction potential in V vs O<sub>2</sub> (i.e., per 1/4 mol of O<sub>2</sub> gas),  $E_g$  is the SCAN+*U* band gap at the  $\Gamma$ -point in eV, and  $E_{hull}$  is the energy above the convex hull in eV/atom. Non-blue markers correspond to qualitative outliers. The dashed black line corresponds to perfect agreement between SCAN+*U* and the model.

work: (1) we have not yet evaluated optimal U values for their oxides within the SCAN+U framework and (2) we wanted to make the construction of our database computationally tractable. For each composition, we considered the following six lattice systems (in Hermann–Mauguin<sup>93</sup> notation) common to perovskites in the Inorganic Crystal Structure Database (ICSD):<sup>94</sup> cubic (e.g., *Pm3m* SrTiO<sub>3</sub>, ICSD #181652), hexagonal (e.g., *P6<sub>3</sub>/mmc* SrMnO<sub>3</sub>, ICSD #185417), rhombohedral (e.g., *R3c* LaCrO<sub>3</sub>, ICSD #167590), tetragonal (e.g., *P4mm* BaTiO<sub>3</sub>, ICSD #164388), orthorhombic (e.g., *Pnma* CaTiO<sub>3</sub>, ICSD #165801), and monoclinic (e.g., *P2<sub>1</sub>/b* CeVO<sub>3</sub>, ICSD #162747). We used the scheme in ref 7 to obtain structures for primitive cells. Figure S1 in the Supporting Information has additional details on the composition of the data set.

**Ab Initio Thermodynamics.** In this work, we calculated neutral  $V_O$  formation energies  $(E_v)$  in the dilute vacancy limit (see Figure S2 for the distribution of  $E_v$  and the data files for tabulated  $E_v$  values). We used the structure\_matcher<sup>95</sup> module in pymatgen<sup>96</sup> to obtain all possible symmetry-distinct neutral  $V_O$  configurations. The formation energy of a neutral  $V_O$  is given by:

$$E_{\rm v} = E_{defective}^{\rm SCAN+U} - E_{pristine}^{\rm SCAN+U} + \frac{1}{2} E_{\rm O_2}^{\rm SCAN}$$
(1)

where  $E^{\text{SCAN}+U}$  is the SCAN+U total energy, pristine is a supercell of the primitive cell, and *defective* is the same supercell with a neutral V<sub>0</sub>.  $E_{O_2}^{\text{SCAN}}$  is the SCAN total energy of an O<sub>2</sub> gas molecule in its groundstate triplet electronic configuration.  $E_v$  is converged to within ±0.1 eV for a 3 × 3 × 3 supercell of the cubic and tetragonal phases, 2 × 2 × 2 supercell of the orthorhombic and monoclinic phases, and 2 × 2 × 1 supercell of the hexagonal and rhombohedral phases (see Figure S3 in the Supporting Information for the convergence of  $E_v$  in different crystal systems with respect to supercell size).

For all perovskites considered, we quantified the stability with  $E_{hull}$ and tested the stability against all elemental phases (i.e., A, B, and O; XC = SCAN) and ordered (i.e., all sites in a given structure have integer atomic occupations), experimentally characterized binary and ternary metal oxides (e.g., AO,  $B_2O_3$ ,  $AB_2O_4$ , etc.; XC = SCAN+U) containing the elements of the perovskite. For the DFT calculations of elemental solids, we used the SCAN XC functional without Hubbard U corrections, as the latter fails in predicting the properties of metallic systems with delocalized electrons. While DFT-SCAN and SCAN+U total energies are incompatible theoretically, this only influences A-rich and B-rich regions of the phase diagram that are isolated from the stability regions of the perovskite compositions we considered and only exist for the sake of completing the individual ternary phase diagrams. Therefore, the  $E_{\rm hull}$  values reported here were calculated using a consistent theoretical treatment of all potential structures using SCAN+U. We used the phase\_diagram module<sup>97,98</sup> in pymatgen<sup>96</sup> to generate the 0 K phase diagrams and calculate  $E_{\rm hull}$  (see Figure S4 in the Supporting Information for the distribution of  $E_{\rm hull}$ ).

To calculate crystal bond dissociation energies (vide infra), we first calculated cohesive energies from atomic total energies (see Figure S5 in the Supporting Information for the periodic trends in experimental cohesive energies). We tested convergence with respect to cell size (12, 16, and 20 Å) and used nearly cubic cells with minimal orthorhombic distortions to lift spurious intrashell degeneracies of atoms with partially filled shells (see Figure S6 in the Supporting Information for the cell-size convergence of atomic total energies). Our predicted magnetic moments agree with the experimental ground-state atomic electron configurations with the exception of V, which prefers [Ar]4s<sup>0</sup>3d<sup>5</sup> over [Ar]4s<sup>2</sup>3d<sup>3</sup> by 0.29 eV—this could be explained by inaccuracies of conventional spin-neutral PAW potentials. $^{99-103}$  With that being said, this only leads to a small change of 0.05 eV in the V-O crystal bond dissociation energies. Finally, we used the Huber regressor in scikit-learn<sup>104</sup> for supervised learning of E<sub>v</sub>. The Huber regressor is a regularized linear regression model that is robust to outliers. While nonlinear regression may find better descriptors for many properties (e.g., symbolic regression identified a simple descriptor for the oxygen evolution reaction activity of metal-oxide perovskites<sup>105</sup>), we find that  $E_v$  can be described by a physically intuitive extended Born-Haber cycle (vide infra), justifying the use of a linear model. The coefficients, intercepts, and scores reported here were cross-validated by averaging over 1000 models trained on random permutations of 50% of the data.

#### RESULTS

**Model for V<sub>o</sub> Formation in Metal-Oxide Perovskites.** First, we report the performance of and key metrics in our best  $E_v$  model for room-temperature-stable ( $E_{hull} \leq 0.025$  eV/atom  $\approx 298.15$ K· $k_B$ , where  $k_B$  is the Boltzmann constant) perovskites, as well as all perovskites considered. Figure 3 shows the agreement between SCAN+U-calculated (vertical axis) and model-predicted (horizontal axis)  $E_v$  values for 142 unique V<sub>o</sub>s

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**Figure 4.** Specific trends in the SCAN+U-calculated and experimental  $E_b$  values vs (a) A-site atoms (+2/+3/+4) oxidation states, (b) B-site atoms (+4), (c) B-site atoms (+3), and (d) B-site atoms (+2). Experimental formation energies/enthalpies are not available for metastable FeO<sub>2</sub>, CoO<sub>2</sub>, and NiO<sub>2</sub>.

in room-temperature-stable perovskites (Figure 3a) and an allencompassing superset of 341 perovskites (Figure 3b), with SCAN+U-calculated and model-predicted values ranging from -2.69 to 9.11 eV and from -0.65 to 7.44 eV, respectively. To the best of our knowledge, this constitutes the most comprehensively trained model for perovskite  $E_v$  values to date. The model is linear with four intuitive energy terms, which we describe in more depth below: crystal bond dissociation energy sum ( $\sum E_b$ ), maximum crystal reduction potential  $(V_r)$ , SCAN+U band gap at the  $\Gamma$ -point  $(E_g)$ , and the energy above the convex hull  $(E_{hull})$ . Note that the crystal system correlates indirectly with  $E_v$  via  $E_g$  and  $E_{hull}$  (see Figure S7 in the Supporting Information). The model's predictions are in good agreement with SCAN+U calculations, with MAEs of  $0.45 \pm 0.04$  and  $0.71 \pm 0.05$  eV for room-temperature-stable structures and all structures, respectively. Figure 3b shows that there are approximately six major qualitative outliers (nonblue markers), which we will address in the Discussion.

**Crystal Feature Engineering Used in the Model.** *Crystal Bond Dissociation Energy.* The inspiration for  $\sum E_b$  stems from its molecular analogue, bond dissociation energies (BDEs), for which experimental values are widely available in references such as the *CRC Handbook of Chemistry and Physics.*<sup>106</sup> For transition-metal (M) oxides, however, experimental BDEs are only available for diatomic molecules/ cations, e.g., MO(g) and MO<sup>+</sup>(g), which lack the inclusion of effects from the metal's oxidation state and the crystal field of the solid. To incorporate these effects, we define a crystal O<sup>2–</sup>–M<sup>n+</sup> bond dissociation energy ( $E_b$ ) as:

$$E_{b}[O^{2-}-M^{n+}] = E_{c}[MO_{n/2}]/N_{b}[O^{2-}-M^{n+}]$$
(2)

where *n* is the oxidation state of M,  $E_c[MO_{n/2}]$  is the cohesive energy of the ground-state polymorph of the binary oxide crystal containing M<sup>*n*+</sup>, and  $N_b[O^{2-}-M^{n+}]$  is the number of  $O^{2-}-M^{n+}$  bonds per  $MO_{n/2}$  formula unit.  $E_b$  can be thought of as the cohesive energy per bond in an ionic oxide framework or the energy required to fully break an  $O^{2-}-M^{n+}$  bond. We use the  $E_c$  values of ground-state binary-oxide polymorphs because they provide a consistent (upper) limit on  $E_b$  and they can be extracted from either experiment or theory.

To calculate  $E_{\rm b}$ , consider the following series of chemical reactions:

$$MO_{n/2}(s) \to M(s) + (n/4)O_2(g), -\Delta E_f[MO_{n/2}]$$
 (3a)

$$M(s) \rightarrow M(g), \qquad E_c[M] \quad (3b)$$

$$(n/4)O_2(g) \to (n/2)O(g),$$
  $(n/4)BDE[O_2]$  (3c)

$$MO_{n/2}(s) \to M(g) + (n/2)O(g)$$
  

$$E_{c}[MO_{n/2}] = -\Delta E_{f}[MO_{n/2}] + E_{c}[M] + (n/4)BDE[O_{2}]$$
(3d)

where  $\Delta E_{\rm f}[{\rm MO}_{n/2}]$  is the binary metal-oxide formation energy/enthalpy,  $E_c[M]$  is the cohesive energy of the pure metal element, and  $BDE[O_2]$  is the bond dissociation energy of  $O_2$  per atom. The energy change for the net reaction (eq 3d) is  $E_c[MO_{n/2}]$ . The experimental  $E_c[MO_{n/2}]$  value can be calculated from the experimental  $\Delta E_f[MO_{n/2}]$ ,<sup>107–111</sup>  $E_c[M]$ ,<sup>112</sup> and BDE[O<sub>2</sub>] values.<sup>106</sup> The experimental  $N_{\rm b}[{\rm O}^{2-}-{\rm M}^{n+}]$  value can be obtained from the experimental binary-oxide crystal structures, which can be found in crystallographic databases such as the ICSD.<sup>94</sup> In the absence of experimental data, theoretical  $E_c[MO_{n/2}]$  and  $N_b[O^{2-}-M^{n+}]$ values can be computed from SCAN+U total energies and optimized crystal structures, which are available in computational databases such as the Materials Project (MP),<sup>113</sup> Open Quantum Materials Database (OQMD),<sup>114,115</sup> and Automatic-FLOW for Materials Discovery (AFLOW).<sup>116</sup> Note that because we define  $E_{\rm b}$  for stable binary metal oxides, namely their ground-state polymorphs, ternary instabilities are captured primarily by  $E_{\text{hull}}$ .

As an example, take the  $E_b[O^{2-}-Ce^{4+}]$  value for  $CeO_2$ , where we are given that M = Ce, n = 4,  $MO_{n/2} = CeO_2$ , and its ground-state polymorph adopts the fluorite structure ( $Fm\overline{3}m$ ). According to the net reaction in eq 3d, the theoretical  $E_c$  value is given by:

where  $E^{SCAN+U}$ [Ce] and  $E^{SCAN}$ [O] are the SCAN+U and DFT-SCAN total energies of gaseous neutral Ce and O in their ground-state  ${}^{1}G_{4}$  and  ${}^{3}P_{2}$  electron configurations, respectively,<sup>117</sup> and  $E^{SCAN+U}[CeO_{2}]$  is that of fluorite CeO<sub>2</sub>. Note that using SCAN+U instead of DFT-SCAN to calculate the total energy of an isolated, neutral Ce atom is appropriate, since Ce is metallic only in its bulk solid ground state and the tightly held open-shell d and f electrons require the Hubbard Ucorrection to mitigate the self-interaction error within DFT. That said, the <sup>1</sup>G<sub>4</sub> atomic state is an open-shell singlet state (coupling a 4f and a 5d electron), requiring two determinants to properly describe its spin coupling. Instead, singledeterminant DFT+U theory only describes a spin-polarized, spin-contaminated singlet. As it turns out, the  $E_c$  error caused by this spin contamination is acceptably small: the theoretical and experimental  $E_{c}[CeO_{2}]$  values are in good agreement (20.51 and 20.78 eV/formula unit, respectively), with an absolute error (AE) of 0.09 eV/atom across all metal oxides considered (see Table S2 in the Supporting Information). Here, the experimental  $E_c$  is given by the formula in eq 3d. Since  $O^{2-}$  is tetrahedrally coordinated to four  $Ce^{4+}$  atoms and there are two  $O^{2-}$  atoms per formula unit of CeO<sub>2</sub>,  $N_{\rm h} = 8$ . Inserting  $E_c$  and  $N_b$  into eq 2 yields theoretical and experimental  $E_b[O^{2^-}-Ce^{4+}]$  values of 2.56 and 2.60 eV/bond, respectively (AE = 0.04 eV). Interestingly, the experimental  $E_{\rm b}[O^{2-}-Ce^{4+}]$  value is 3.15–3.40 times smaller than its molecular analogues, BDE[CeO(g)] (8.19 eV at 298 K) and BDE[CeO<sup>+</sup>(g)] (8.83  $\pm$  0.16 eV at 298 K).<sup>106</sup> Ce in  $CeO_2(s)$  donates its four valence electrons  $(6s^25d^14f^1)$  equally among its nearest O neighbors, within the electrostatic potential of the CeO<sub>2</sub> crystal, whereas Ce in CeO(g) and  $CeO^{+}(g)$  shares its electrons with only one O absent any other potential, creating up to a triple M-O bond in the diatomic species.<sup>118</sup> Additionally, differences in the ionic characters of the  $O^{2-}-Ce^{4+}$ , O-Ce (i.e.,  $\approx O^{2-}-Ce^{2+}$ ), and  $O-Ce^+$  (i.e.,  $\approx O^{2-}-Ce^{3+}$ ) bonds in  $CeO_2(s)$ , CeO(g), and  $CeO^+(g)$ , respectively, also contribute to the deviation between molecular and crystal BDEs.

Figure 4 shows the specific trends in the theoretical  $E_{\rm b}$  values (vertical axis) for the ground-state  $MO_{n/2}$  (horizontal axis) of the A- and B-site cations that we considered. The experimental (unfilled markers) and theoretical (filled markers)  $E_{\rm b}$  values are in good agreement, with MAE = 0.04 eV, indicating that the former can be used when the latter are unavailable (or vice versa), without loss of accuracy. For the A-site cations (Figure 4a), the theoretical  $E_{\rm b}$  trend from strong to weak crystal bonds is  $Ce^{4+}$  (2.56 eV) >  $La^{3+}$  (2.50 eV) >  $Ce^{3+}$  (2.48 eV) >  $Ca^{2+}$  $(1.88 \text{ eV}) > \text{Sr}^{2+} (1.76 \text{ eV}) > \text{Ba}^{2+} (1.70 \text{ eV})$ . The higher  $E_{\text{h}}$ values for Ce and La in comparison to those for the alkaline earths can be attributed to either (1) the higher oxidation states of Ce (+4 or +3) and La (+3), which can increase the magnitude of the electrostatic potential energy associated with each bond, or (2) the greater number of electrons per bond for  $CeO_2$  (0.5 e<sup>-</sup>/bond) in comparison to  $La_2O_3/Ce_2O_3$ (0.43 e<sup>-</sup>/bond) and CaO/SrO/BaO (0.33 e<sup>-</sup>/bond). Additionally, the increasing size mismatch between  $A^{2+}$  and  $O^{2-}$ from Ca<sup>2+</sup> to Ba<sup>2+</sup> results in a weaker electrostatic attraction, leading to weaker crystal bonds, thus explaining the observed trend in  $E_{\rm b}$ .

For the  $B^{4+}$  cations (Figure 4b), the theoretical  $E_{\rm b}$  values display a monotonically decreasing trend as B progresses from Ti to Ni: i.e.,  $Ti^{4+}$  (3.16 eV) >  $V^{4+}$  (2.90 eV) >  $Cr^{4+}$  (2.61 eV)  $> Mn^{4+} (2.25 \text{ eV}) > Fe^{4+} (2.13 \text{ eV}) > Co^{4+} (2.01 \text{ eV}) > Ni^{4+}$ (1.96 eV). We find a similar trend for the  $B^{3+}$  cations (Figure 4c), i.e.,  $Ti^{3+}$  (2.62 eV) >  $V^{3+}$  (2.53 eV) >  $Cr^{3+}$  (2.32 eV) >  $Fe^{3+} = Co^{8/3+}$  (2.07 eV) >  $Mn^{3+}$  (1.97 eV) >  $Ni^{3+}$  (1.81 eV), with Fe and Co as notable exceptions. The  $B^{2+}-O^{2-}E_b$  values (Figure 4d) are lower than the corresponding  $B^{3+}-O^{2-}E_{h}$ values, which in turn are consistently lower than the respective  $B^{4+}-O^{2-}E_{h}$  values (except for Co). Notably, Fe<sup>3+</sup> is stabilized by its  $d^5$  electron configuration, leading to a higher  $E_{\rm b}$  value in comparison to  $Mn_2O_3$ . On the other hand, the  $E_{\rm h}$  value for  $Co^{8/3+}$ , which is derived from  $Co(II,III)_3O_4$  and acts as a proxy for the unstable  $Co(III)_2O_3$ , is higher than that for  $CoO_2$ , which can be attributed to the stability (metastability) of  $Co_3O_4$  (CoO<sub>2</sub>). Figure S8 in the Supporting Information has additional details on  $E_{\rm b}$  and its molecular analogues.

Crystal Reduction Potential. In the same spirit as  $\sum E_{b}$ ,  $V_r$  extends useful concepts from atomic/molecular (electro)chemistry, such as ionization energies  $(E_i)$  and standard reduction potentials  $(V^\circ)$ , to the solid state by consistently including crystal field effects and removing any solvation effects in an aqueous (or other liquid) medium. Another motivation for  $V_r$  stems from the unavailability of experimental  $V^\circ$  values for the following reduction reactions, relevant in ABO<sub>3</sub> perovskites: Fe<sup>4+</sup>  $\rightarrow$  Fe<sup>3+</sup> and Ni<sup>4+</sup>  $\rightarrow$  Ni<sup>3+ 106</sup> We define a crystal M<sup>n+</sup>  $\rightarrow$  M<sup>m+</sup> (where n > m) reduction potential  $(V_r)$  vs O<sub>2</sub> (per 1/4 mol of O<sub>2</sub> gas) as:

$$V_{\rm r}[{\rm M}^{n+} \to {\rm M}^{m+}] = -E_{\rm r}[{\rm M}^{n+} \to {\rm M}^{m+}]/(n-m)F$$
 (5)

where *n* and *m* are the oxidation states of the oxidized and reduced metals in the ground-state polymorphs of their binary metal-oxide crystals  $MO_{n/2}$  and  $MO_{m/2}$ , respectively, *F* is the Faraday constant, and:

$$E_{\rm r}[{\rm M}^{n+} \rightarrow {\rm M}^{m+}]$$

$$= E^{{\rm SCAN}+U}[{\rm MO}_{m/2}] + (n-m)E^{{\rm SCAN}}[{\rm O}_2]/4$$

$$- E^{{\rm SCAN}+U}[{\rm MO}_{n/2}] \qquad (6)$$

Note that  $V_r$  in eq 5 is strictly defined for the Gibbs energy of reduction, which we approximate here as the internal energy (i.e.,  $G \approx E$ ), ignoring p-V and entropic effects, in order to apply our scheme readily to large data sets.

 $V_r$  can be interpreted as the reduction potential of a metal cation in a crystal. For example, consider  $V_r$ [Ce<sup>4+</sup>  $\rightarrow$  Ce<sup>3+</sup>], where M = Ce, n = 4, m = 3, n - m = 1, MO<sub>n/2</sub> = CeO<sub>2</sub> (ground-state polymorph = fluorite structure, *Fm*3*m*), and MO<sub>m/2</sub> = CeO<sub>3/2</sub> =  $^1/_2$ Ce<sub>2</sub>O<sub>3</sub> (ground-state polymorph = hexagonal lattice, *P*3*m*1). Following from eq 6, the theoretical  $E_r$  value is:

$$E_{\rm r}[{\rm Ce}^{4+} \to {\rm Ce}^{3+}] = \frac{1}{2} E^{{\rm SCAN}+U}[{\rm Ce}_2{\rm O}_3] + E^{{\rm SCAN}}[{\rm O}_2]/4$$
$$- E^{{\rm SCAN}+U}[{\rm CeO}_2] = 1.82 \text{ eV}$$
(7)

On the other hand, the experimental  $E_r = \frac{1}{2}\Delta E_f[Ce_2O_3] - \Delta E_f[CeO_2] = 1.86 \text{ eV.}^{77}$  Inserting  $E_r$ , n - m = 1, and F into eq 5 yields theoretical and experimental  $V_r$  values of -1.82 and -1.86 V per 1/4 mol of O<sub>2</sub>, respectively (AE = 0.04 V per 1/4 mol of O<sub>2</sub>). Thus, reducing Ce<sup>4+</sup> to Ce<sup>3+</sup> in its bulk oxide state



**Figure 5.** Comparison of experimental (blue) and theoretical (orange)  $V_r$  values with respect to available experimental aqueous standard reduction potentials (green),<sup>106</sup> showing that the qualitative trends are obeyed. Experimental  $V^{\circ}$  values are not available for Fe<sup>4+/3+</sup>, Ni<sup>4+/3+</sup>, and Ni<sup>3+/2+</sup>.



**Figure 6.** Relationship between SCAN+U-predicted  $E_v$  and  $\sum E_b$  (blue),  $V_r$  (orange),  $E_g$  (green), and  $E_{hull}$  (red). The numerical ranges of the first three descriptors (i.e., the difference between the largest and smallest horizontal axis values) are approximately equal (i.e., ~5 eV or V).  $V_r$  captures most of the contribution to  $E_v$ .

should theoretically (experimentally) require a reduction potential of 1.82 V (1.86 V) per 1/4 mol of oxygen gas.

Figure 5 shows that the experimental (blue bars) and theoretical (orange bars)  $V_r$  values are in good agreement, with MAE = 0.10 V vs  $O_2$ , implying that, like  $E_b$ , experimental values can be used to make accurate predictions in the absence of theoretical data (and vice versa). The aqueous  $V^{\circ}$  values (green bars), however, poorly capture the qualitative trends in  $V_r$ . Note that we plot  $V^{\circ}$  with respect to the standard hydrogen electrode, whereas both theoretical and experimental  $V_r$  values are plotted with respect to 1/4 mol of  $O_2(g)$ . While the V°s of the transition-metal cations (Figure 5b-d) capture most of the trends in  $V_r$  (but have the wrong sign for V<sup>4+</sup>, Cr<sup>4+</sup>, Mn<sup>4+</sup>, Mn<sup>3+</sup>, Fe<sup>3+</sup>, and Co<sup>3+</sup>), those of the alkaline-earth-metal and lanthanide cations (Figure 5a) do not. Hence, it is critical to use  $V_{\rm r}$  and not  $V^{\circ}$  values because the latter lack the consistent inclusion of crystal field effects required for a quantitatively accurate description of solid-state cation reduction. For specific trends in  $V_r$ , please see Figure S9 in the Supporting Information.

**Model Intuition.** Having presented  $E_b$  and  $V_r$  as descriptors for  $E_v$  and clarified how they relate to fundamental chemical concepts and principles, we now inspect the model's

construction and offer intuitive rationalizations for the signs and magnitudes of its coefficients (c). Our model consists of four terms (Figure 3),  $\sum E_b$  representing the energy penalty of breaking bonds and eventually creating an O2 molecule from an  $O^{2-}$  ion (*O*-*M* bond dissociation in Figure 1),  $V_r$  quantifying the energy required to reduce "adjacent" cations due to the electrons left behind by the removed  $O^{2-}$  (M reduction in Figure 1),  $E_{g}$  signifying the band structure cost of placing additional electrons in an oxygen-deficient perovskite in its conduction band (not shown in Figure 1 but affects M *reduction*), and  $E_{\text{hull}}$  highlighting the impact of the underlying (in)stability of the structure in controlling V<sub>O</sub> formation (metal oxide stability in Figure 1). Our model, with energy terms related to atomization  $(\sum E_{\rm b})$ , ionization  $(V_{\rm r})$ , and polarity of chemical bonds  $(E_g)$ , can be considered to be an extended Born-Haber cycle.

The first term,  $\sum E_b$ , is the sum of the  $E_b$  values for the crystal bonds formed between a specific  $O^{2-}$  that forms the vacancy and its nearest cation neighbors. For perovskites:

$$\sum E_{\rm b} = 4E_{\rm b}[{\rm O}^{2-}-{\rm A}^{n+}] + 2E_{\rm b}[{\rm O}^{2-}-{\rm B}^{m+}]$$
(8)



because there are four  $O^{2-}-A^{n+}$  and two  $O^{2-}-B^{n+}$  crystal bonds per  $O^{2-}$ . Note that our definition of  $\sum E_b$  is quite general and easily extends to any desired metal-oxide material. If The sign of the  $E_b$  coefficient,  $c_b > 0$ , is consistent with the intuition that energy is required to break crystal bonds (R =0.77, blue markers in Figure 6). The magnitude of  $c_b$  is smaller (0.1-0.2) than those of the other coefficients (0.3-55.8,  $c_b <$  $c_g < |c_r| < |c_{hull}|$ ), suggesting that  $\sum E_b$  serves primarily as a correction factor to other terms in the model. Differences between the bond orders in binary and ternary oxides also can affect the magnitude of  $c_b$ ; quantifying such effects will be the

focus of future work. The second term,  $V_r$ , is the maximum  $V_r$  value among the nearest cation neighbors of a specific  $O^{2-}$  and is the most dominant factor determining  $E_v$  in our model. Indeed,  $E_v$  anticorrelates quite strongly with  $V_r$  (orange markers in Figure 6), with R = -0.88. Choosing the maximum  $V_r$ , we assume that the most reducible nearest cation neighbor undergoes reduction and hence governs the  $V_r$  contribution to  $E_v$ . For perovskites:

$$V_{\rm r} = \max(V_{\rm r}[{\rm A}^{m^+} \to {\rm A}^{n^+}], V_{\rm r}[{\rm B}^{r^+} \to {\rm B}^{s^+}])$$
(9)

where m > n and r > s. The sign (<0) and magnitude (1.5–1.6) of the  $V_r$  coefficient ( $c_r$ ) resembles the energy for an *N*-electron reduction reaction,  $E_r = -NV_r$ , where N = 2, N < 2, and  $N \ll 2$  for  $V_O$  formation in ionic, ionic + covalent, and metallic oxides, respectively. Therefore,  $-c_r$  can be interpreted physically as the number of electrons donated by a specific  $O^{2-}$  to its most reducible nearest cation neighbors upon  $V_O$  formation in an ionic + covalent metal oxide.

Electrons from the departing oxygen that are left behind occupy the conduction band rather than the valence band. Hence the band gap qualitatively describes the cost of adding electrons to the conduction band upon reduction, which results in the coefficient of the third term,  $E_g$ ,  $c_g > 0$  (R = 0.34, green markers in Figure 6). Note that this gap is the one at the  $\Gamma$ -point obtained from a fully self-consistent SCAN+Ucalculation. Typical band structure and density of states calculations involve a non-self-consistent calculation after the completion of a fully self-consistent calculation. However, to minimize computational costs, we have not performed a nonself-consistent calculation, which is sufficient for our purposes here. Since the numerical ranges of the first three descriptors are approximately equal, the magnitude of  $c_{\rm g}$  (0.3–0.4) signifies that  $E_{\rm g}$  has a 15–20% weight to the overall  $E_{\rm v}$ .

Finally, the  $\vec{E}_{hull}$  coefficient,  $c_{hull} < 0$  (R = -0.18, red markers in Figure 6), reflects the intuition that instability favors decomposition: e.g., via facile  $V_{\Omega}$  formation. Note that the  $c_{\text{hull}}$ coefficients do vary significantly (5.5-55.8) between our models trained on room-temperature-stable and all perovskites in our data set, likely offsetting the actual  $E_{\rm hull}$  values that are used in the room-temperature-stable ( $E_{\text{hull}} \leq 25 \text{ meV/atom}$ ) and the full (no upper limit on  $E_{hull}$ ) perovskite sets. In both of our theoretical models, chull exhibits the highest magnitude of the coefficients; however, the typical values of  $E_{\text{hull}}$  for crystals are in the range of 0-100 meV/atom, which is at least one order of magnitude lower than  $E_{\rm b}$ ,  $V_{\rm r}$ , or  $E_{\rm g}$ , resulting in a significantly smaller contribution for those structures that are stable ( $E_{\text{hull}} = 0$ ) and mildly unstable ( $E_{\text{hull}} \approx 50 \text{ meV/atom}$ ). Importantly, ours is the first modeling work to include the effects of thermodynamic (in)stability on Vo formation via  $E_{\text{hull}}$ , which could be a source of the inaccuracy and/or nonintuitiveness of previous models.<sup>33,51-69</sup>

B-Site Trends for Materials Design and Candidate Identification. In this section, we discuss the specific trends in the SCAN+U-calculated and model-predicted  $E_v$  with respect to changes in B and its oxidation state (n), focusing on materials design and candidate identification for various applications. Figure 7 shows the SCAN+U-calculated  $E_v$ distribution (DFT+U, circles), average SCAN+U-calculated  $E_v$  values ((DFT+U), filled diamonds), and average modelpredicted  $E_v$  values ((Model), unfilled diamonds, vertical axis) as a function of B (horizontal axis) and n (panels). Overall, the average SCAN+U-calculated and model-predicted  $\langle E_v \rangle$  values are in good agreement (MAE = 0.45 eV). For n = 4 (i.e., B<sup>4+</sup>),  $\langle E_v \rangle$  decreases in the order Ti > V > Mn  $\approx$  Cr > Ni > Co  $\approx$  Fe (Figure 7a), which largely coincides with increasing  $V_r[B^{4+} \rightarrow$  $B^{3+}$ ] (R = -0.98): namely, Ti (-1.91 V vs O<sub>2</sub>) < V (-0.92 V vs  $O_2$ ) < Cr (-0.42 V vs  $O_2$ )  $\approx$  Mn (-0.40 V vs  $O_2$ ) < Ni  $(0.40 \text{ V vs } O_2) < \text{Co} (0.55 \text{ V vs } O_2) < \text{Fe} (0.97 \text{ V vs } O_2).$ 

Α	В	$\sum E_{b}$	$V_{ m r}$	$E_{g}$	space group	model $E_{\rm v}$ (eV)	$T_{\rm m}$ (K)
Y <sup>3+</sup>	Fe <sup>3+</sup>	16.23	-1.45	2.43 <sup>131</sup> (exptl)	Pnma	4.76	1993 <sup>132</sup>
Y <sup>3+</sup>	Co <sup>3+</sup>	16.23	-1.02	$1.48^{133}$ (theor)	Pnma	4.34	n/a
Bi <sup>3+</sup>	Fe <sup>3+</sup>	10.71	-0.99	2.50 <sup>134</sup> (exptl)	R3c	3.99	1234-1336 <sup>135</sup>
Bi <sup>3+</sup>	Co <sup>3+</sup>	10.71	-0.99	1.70 <sup>136</sup> (exptl)	P4mm	3.80	729 <sup>126</sup>

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 Table 1. List of Promising Perovskites for STCH<sup>a</sup>

<sup>*a*</sup>Compounds in boldface type might exhibit simultaneous reduction of the A- and B-site cations. The model  $E_v$  value is calculated from the experimental  $\sum E_b$  value (using eq 8), experimental  $V_r$  value (using eq 9), experimental (exptl) or theoretical (theor)  $E_g$  value, and MP PBE+U-calculated  $E_{hull}$  value (not listed because all  $E_{hull} \approx 0$ ). We include the measured melting temperatures ( $T_m$ ) of the candidates because thermal stability is also a significant screening criterion for STCH.

Similarly, for n = 3 (i.e.,  $B^{3+}$ ), the  $\langle E_v \rangle$  trend (Figure 7b) nearly exactly anticorrelates with that of  $V_r[B^{3+} \rightarrow B^{2+}]$  (R = -1.00): Cr (-2.87 V vs O<sub>2</sub>) < Ti (-2.75 V vs O<sub>2</sub>) < V (-2.01 V vs O<sub>2</sub>) < Fe (-1.44 V vs O<sub>2</sub>) < Mn (-1.02 V vs O<sub>2</sub>) < Co (-0.90 V vs O<sub>2</sub>) < Ni (0.05 V vs O<sub>2</sub>). Finally, for Co and Ni, Figure 7a-c shows that  $E_v$  increases with decreasing *n*, which is consistent with the relative instability of their higher oxidation states.

For simplicity in the discussion of materials design strategies, we bin  $E_v$  into three discrete ranges: low  $E_v$  ( $E_v \leq 2.5$  eV), intermediate  $E_v$  (2.5 <  $E_v \leq 5$  eV), and high  $E_v$  ( $E_v > 5$  eV). Low- $E_v$  oxides are more ideal for SOFCs, thermochemical O<sub>2</sub> storage and pumping/separation, thermochemical energy storage, and thermochemical air separation, with optimal Bsite cations being Fe<sup>4+</sup>, Co<sup>4+</sup>, and Ni<sup>4+</sup>. While perovskites containing these cations are consistent with state-of-the-art (La,Sr)(Co,Fe)O<sub>3</sub> SOFC cathodes,<sup>60</sup> this reprediction of their optimality highlights the model's accuracy and potential transferability to superternary perovskite metal oxides.

Intermediate- $E_v$  metal oxides are more suitable for STCH (WS and/or CDS) because the interactions between  $O^{2-}$  and its nearest cation neighbors should be neither too strong for extensive thermal reduction nor too weak for spontaneous water splitting. The best B-site cations for STCH are V<sup>4+</sup>, Mn<sup>3+</sup>, Fe<sup>3+</sup>, Co<sup>3+</sup>, Co<sup>2+</sup>, and Ni<sup>2+</sup>. Mn<sup>3+</sup> appears in Ca-, Ce-, and Mn-based ABO3 perovskites, which we recently identified to be promising candidates that potentially exhibit simultaneous reduction of both Ce4+ (A-site) and Mn3+ (B-site).7 While the model-predicted  $E_v$  value for Mn<sup>4+</sup> seems like it is too low for STCH,  $BaCe_{1/4}Mn_{3/4}O_3$  is one of the best water splitters to date.<sup>119</sup> This underestimation is likely due to differences between its  $E_g$  value and that of BaMnO<sub>3</sub> (since Mn<sup>4+</sup> is the redox-active cation in both stable materials and therefore their  $\sum E_{\rm b}$ ,  $V_{\rm r}$ , and  $E_{\rm hull}$  values should be similar), thus making band gap prediction for superternary perovskite metal oxides, an important avenue for improving the model. On the basis of these B-site cations, there are three ternary prototypes worth exploring further: (1)  $A^{2+}V^{4+}O_{3}$ , (2)  $A^{3+}(Mn,Fe,Co)^{3+}O_3$ , and (3)  $A^{4+}(Co,Ni)O_3$ . For prototype 1, while the phase impurity of oxides containing  $V^{120-122}$ reduces the likelihood of its practical use,  $P2_1/c$  SrVO<sub>3</sub> is a reasonable candidate with a SCAN+U-predicted  $E_v$  value of 3.75-3.78 eV.

For prototype 2 or 3, we used the following scheme for enumerating promising candidates, outside of our training data. First, we screened for elements known to exist most commonly in the 3+/4+ oxidation state.<sup>123</sup> Second, we selected only those 3+/4+ cations whose ionic radii ( $r_A$ ) produce a Goldschmidt tolerance factor<sup>91</sup> (t) greater than 0.71 (i.e., the stability limit of the perovskite structure), where:

$$t = \frac{r_{\rm A} + r_{\rm B}}{\sqrt{2} (r_{\rm B} + r_{\rm O})}$$
(10)

Article

 $r_{\rm B}$  is the ionic radius of the six-fold-coordinated B<sup>3+/2+</sup>, and  $r_{\rm O}$ is the ionic radius of the six-fold-coordinated  $O^{2-}$  (1.40 Å). Note that t has a positivity rate of 74%, so even if the tolerance criterion predicts a perovskite structure to form, the composition still may not actually form. Recently, Bartel et al. developed a more accurate tolerance factor (positivity rate of 92%);<sup>124</sup> however, we use eq 10 because its solution is analytical and therefore easier to solve for the relevant range of  $r_{\rm B}$  for which t > 0.71. That said, a numerical solution of the new tolerance factor<sup>124</sup> for  $r_{\rm B}$  is not difficult to obtain and can be a strategy for subsequent studies. We selected only those compositions whose MP-PBE+U-calculated  $E_{hull} \leq$ 0.025 eV/atom: i.e., those that are predicted to be stable at room temperature. Table 1 gives the particularly promising candidates. (La,Ce)<sup>3+</sup>(Mn,Fe,Co)<sup>3+</sup>O<sub>3</sub> have been screened theoretically as candidates before by Emery et al.,<sup>39</sup> whereas those in Table 1 are new materials with known synthesis procedures. For example,  $Bi(Fe,Co)O_3$  has a model  $E_v$  value in the target range of  $\approx 3.4 - 3.9 (\pm 0.1)$  eV for improved STCH materials.' Of these, BiFeO<sub>3</sub> is especially interesting because it is a promising multiferroic material with several synthesis routes already proven.<sup>125</sup> In addition, the absolute differences between the Bi<sup>3+</sup> and Fe<sup>3+</sup>  $V_r$  values ( $|V_r[Bi^{3+} \rightarrow Bi^0] - V_r[Fe^{3+} \rightarrow Fe^{2+}]| = 0.46 \text{ V vs } O_2$ ) and the Bi<sup>3+</sup> and Co<sup>8/3+</sup>  $V_r$  values ( $|V_r[Bi^{3+} \rightarrow Bi^0] - V_r[Co^{8/3+} \rightarrow Co^{2+}]| = 0.03 \text{ V vs } O_2$ ) are small in comparison to that of the simultaneously redox-active Ce4+ and Mn3+ cations in Ca-, Ce-, and Mn-based ABO3 perovskites  $(|V_r[Ce^{4+} \rightarrow Ce^{3+}] - V_r[Mn^{3+} \rightarrow Mn^{2+}]| = 1.00 \text{ V}$ vs  $O_2$ ).<sup>7</sup> Consequently, Bi(Fe,Co)O<sub>3</sub> might exhibit higher simultaneous redox activity and therefore a larger entropy of reduction. On the negative side, BiFeO<sub>3</sub> melts at a temperature  $(T_{\rm m})$  of 1234–1336 K, which is lower than a STCH cycle demands ( $\approx$ 2000 K). Encouragingly, the  $T_{\rm m}$  values can be increased via Co doping,<sup>126</sup> so the quaternary species could prove sufficiently robust. Note that we list YFeO<sub>3</sub> as promising—despite its model  $E_v$  (4.76 eV) lying outside the target range ( $\approx$ 3.4–3.9 eV)—because its  $T_{\rm m}$  value (1993 K) is greater than that of the  $E_v$ -ideal Bi(Fe,Co)O<sub>3</sub> (1234–1336 K) and therefore Y-doped Bi(Fe,Co)O<sub>3</sub> might be a promising candidate for STCH.

Finally, high- $E_v$  metal oxides are optimal for semiconductors and ferroelectrics because their performance can be limited by the formation of deleterious defects. For example, V<sub>o</sub>s induce ferroelectric domain wall pinning in BiFeO<sub>3</sub><sup>127</sup> (intermediate  $E_v$ ) but not in A<sup>2+</sup>Ti<sup>4+</sup>O<sub>3</sub><sup>36</sup> (high  $E_v$ ), which restricts polarization switching in the former. The most favorable B-site cations for high  $E_v$  include Ti<sup>4+</sup>, Ti<sup>3+</sup>, V<sup>3+</sup>, and Cr<sup>3+</sup>, which is consistent with the common ferroelectric crystals  $BaTiO_3$  and  $PbTiO_3^{\ 128}$  and those less explored, such as  $CeCrO_3^{\ 129}$  and  $ScCrO_3^{\ 130}$ 

**Model Simplification.** When quantum mechanics calculations are unavailable,  $E_{\rm b}$  and  $V_{\rm r}$  can be calculated from experimental<sup>107–110</sup> formation energies/enthalpies and element cohesive energies.<sup>112</sup> Moreover,  $E_{\rm g}$  can be obtained from the experimental literature or computational databases<sup>113,137–140</sup> or by performing a simpler single-step self-consistent SCAN+U calculation. In cases where  $E_{\rm g}$  is unknown or unreliable, we constructed simplified models for room-temperature-stable perovskites:

$$E_{\rm v} = -1.8V_{\rm r} - 66.0E_{\rm hull} + 2.7\tag{11}$$

and all perovskites:

$$E_{\rm v} = 0.1 \sum E_{\rm b} - 1.7V_{\rm r} - 6.5E_{\rm hull} + 0.5 \tag{12}$$

with MAEs of  $0.50 \pm 0.05$  and  $0.74 \pm 0.05$  eV, respectively. Note that in eq 11 the estimated coefficient for  $\sum E_b$  is  $-0.03 \pm 0.15$  eV and therefore fails to reject the null hypothesis. Figure S10 in the Supporting Information shows the performance of our simplified  $E_v$  models (eqs 11 and 12).

In scenarios where  $E_{\rm hull}$  is undetermined or untrustworthy (e.g., for hypothetical and/or strongly correlated phases that are not present in computational materials databases such as MP, OQMD, and AFLOW), we suggest the following strategy when it is impractical to compute the phase diagram. First, one should use the model for stable perovskites in either Figure 3a (with  $E_{\rm g}$ ) or eq 11 (without  $E_{\rm g}$ ). If  $E_{\rm hull} = 0.025$  eV/atom, then  $c_{\rm hull}E_{\rm hull} = -1.39 \pm 0.40$  and  $-1.65 \pm 0.29$  eV for the model with and without  $E_{\rm g}$ , respectively. Therefore, if the perovskite is room-temperature-stable, then the maximum decrease in  $E_{\rm v}$  is (1.39–1.65)  $\pm$  0.40 eV. Second, since stability often is desired, if the application-specific ideal- $E_{\rm v}$  range encompasses the model  $E_{\rm v}$  value and the model  $E_{\rm v} - 1.39$  value (with  $E_{\rm g}$ ) or  $E_{\rm v} - 1.65$  value (without  $E_{\rm g}$ ), then the material is worth considering.

#### DISCUSSION

**Comparison with a Prevailing Model.** In the first part of the discussion, we compare our model to that of Deml et al.<sup>66</sup> (the state-of-the-art), which was trained on 45 insulating metal oxides, including binary and ternary main-group and transition-metal oxides in the antifluorite, corundum, rock salt, rutile, perovskite, and spinel structures. Note that both Deml et al. and we considered unstable metal oxides in our databases— 58% (42%) of the metal oxides they (we) considered are room-temperature stable. Their model is:

$$E_{\rm v} = 0.72 \left[ |\Delta H_{\rm f}| + 0.60 \left( E_{\rm Op} + \frac{c_{\rm g}}{2} E_{\rm g} + 2.60 \langle \Delta \chi \rangle \right) \right] - 2.07$$
(13)

where  $\Delta H_{\rm f}$  is the metal oxide formation enthalpy in eV/atom,  $E_{\rm Op}$  is the O 2*p* band center in eV,  $\langle \Delta \chi \rangle$  is the average Pauling electronegativity difference between O<sup>2-</sup> and its nearest cation neighbors, and  $c_{\rm g} = 1.00$  (1.50) for the GW-calculated (PBE+U-calculated)  $E_{\rm g}$  in eV. Equation 13 has training and testing MAEs of 0.19–0.21 eV (the first value is for GW and the second is for PBE+U gaps) and 0.39 eV, respectively, where the latter is slightly better than our cross-validated MAE for room-temperature-stable materials (0.45 ± 0.04 eV). When it is normalized by the range of  $E_{\rm v}$ , however, their testing MAE (9%) is slightly worse than ours (6–7%). Despite the differences between some of the terms in our models, there is an underlying relationship between  $\Delta H_{\rm f}$  and  $V_{\rm r}$ . To this end, consider the decomposition of binary  $M^{n+}$  oxides:

$$\mathrm{MO}_{n/2} \to M + (n/4)O_2 \tag{14}$$

where the energy release upon decomposition is:

$$\Delta E_{\rm d} = E_{\rm r}[M^{n+} \to M^0] = -\Delta H_{\rm f} \tag{15}$$

Inserting  $E_r[M^{n+} \rightarrow M^0]$ , n = 4, m = 0, and F into eq 5 yields:

$$V_{\rm r}[{\rm M}^{n+} \to {\rm M}^0] = \frac{\Delta H_{\rm f}}{nF}$$
(16)

Despite the direct mathematical relationship between  $\Delta H_{\rm f}$  and  $V_{\rm r}$ , two important effects are better accounted for with the crystal  $V_{\rm r}$  value used in our model than with  $\Delta H_{\rm f}$ . First, a neutral  $V_{\rm O}$  reduces its neighboring cations; hence, the formation energy per reducing electron  $(V_{\rm r})$  provides a more consistent comparison across metal oxides with varied oxidation states. Second,  $\Delta H_{\rm f}$  neglects the presence of other stable nonzero oxidation states for the reduced metal, while the  $V_{\rm r}$  value defined in our work better accounts for oxidation state changes. For example, Ti<sup>4+</sup>  $\rightarrow$  Ti<sup>3+</sup> is more sensible than Ti<sup>4+</sup>  $\rightarrow$  Ti<sup>0</sup> when CaTiO<sub>3</sub> is undergoing reduction because the  $V_{\rm O}$ -induced two-electron reduction typically involves two or more cation acceptors rather than reduction of one cation all the way to its metallic state.

To provide a more quantitative comparison between our model and that of Deml et al.,<sup>66</sup> we fit a model (using training methods similar to those described in the Computational Methods) to their PBE+U-calculated  $E_v$  values with our descriptors. Note that HgTiO<sub>3</sub> was excluded from the reported model of Deml et al. and its inclusion increases the MAE from the reported 0.21 to 0.27 eV. Our model for their data, i.e.,

$$E_{\rm v} = -1.2V_{\rm r} + 0.3E_{\rm g} + 1.87 \,{\rm eV}$$
 (17)

highlights the important role of  $V_r$  in governing  $E_v$  trends, as it is one of only two metrics required to achieve MAE = 0.25 eV (see Figure 8). Notably, the coefficient we obtain for the  $E_g$ term (0.3) is similar to their model (0.22 or 0.32 depending on whether the PBE+U- or GW-based model is used), while the contribution of other descriptors used in their model drops down to a single  $V_r$  in ours. Additionally, our model is able to reproduce their PBE+U calculated  $E_v$  with lower errors and fewer parameters, highlighting that our descriptors are both more accurate and yield better physical insight.

Revisiting HgTiO<sub>3</sub>, the authors in ref 66 hypothesize that its unpredictability (see the deviation between the orange marker with black outline and the dashed black line) "arises from the inadequate treatment of the filled Hg *d* orbitals in DFT". Our results show, however, that the inclusion of  $V_r$  in the model enables a better prediction (AE = 0.45 eV vs 2.10 eV for our model vs their model, respectively) of the PBE+*U*-calculated  $E_v$  value for HgTiO<sub>3</sub> (3.05 eV). This improvement can be ascribed to the greater reducibility and therefore control over the  $E_v$  value of Hg<sup>2+</sup> (MP-PBE+*U*-calculated  $V_r$ [Hg<sup>2+</sup>  $\rightarrow$  Hg<sup>0</sup>] = -0.65 V vs O<sub>2</sub>) relative to Ti<sup>4+</sup> (MP-PBE+*U*-calculated  $V_r$ [Ti<sup>4+</sup>  $\rightarrow$  Ti<sup>3+</sup>] = -2.25 V vs O<sub>2</sub>), which is captured by  $V_r$  but not by a combination of  $\Delta H_{fr} E_{Op}, E_{gr}$  and  $\Delta \chi$ .

Avenues for Improving and Extending the Model. In the second part of the discussion, we first analyze the six outliers in Figure 3b. On the basis of our low-, intermediate-,



Figure 8. Comparison of our model with the prevailing model in the literature.  $^{66}$ 

and high- $E_v$  bins, we classify these outliers as either quantitative (where the SCAN+U-calculated and modelpredicted  $E_v$  values are in the same bin) or qualitative (where these  $E_v$  values are in different bins). The quantitative outliers are monoclinic (m-)LaCrO<sub>3</sub> ( $P2_1/c$ ) and hexagonal (h-)CaTiO<sub>3</sub> ( $P6_3/mmc$ ), whereas the qualitative outliers are rhombohedral (r-)BaCrO<sub>3</sub> (R3c), tetragonal (t-)LaCoO<sub>3</sub> (P4mm), cubic (c-)LaCoO<sub>3</sub> (Pm3m, ICSD #28921), and c-LaTiO<sub>3</sub> (*Pm3m*, ICSD #28908). In general, multiple possible reasons exist for deviations in the SCAN+U-calculated and model-predicted  $E_v$  values. First, the intrinsic instability ( $E_{hull}$  > 0.042 eV/atom for all six outliers) may catalyze nonlocal changes in the structure that models with local features cannot capture. That said, given the small number of outliers,  $E_{\rm hull}$ captures most of the stability-related contributions to  $E_{\rm v}$ . Second, for simplicity in the screening process, we aligned ferromagnetically the magnetic moments of all cations. Despite this approximation leading to a good description of many basic properties of standard ferromagnetic materials, such as permanent magnetic moments and phase diagrams,141 antiferromagnetism, relevant for the Cr-containing outliers, may lead to inconsistencies between the initial and final spin configurations. Finally, although we considered the high-, intermediate-, and low-spin states for the Co-containing outliers, a more comprehensive analysis of their magnetic structures is left for future work.

Next, we outline avenues for improving the model. One limitation of the model is using  $E_b$  from binary systems, as the actual bond lengths in ternary perovskites (and beyond) structures may be significantly different from the binary oxide bond lengths. So, to account for this, one could either do more calculations to see how  $E_b$  changes with bond length for individual systems or fit a simple function (e.g., an exponent or a Gaussian centered around the binary metal oxide bond length) as a weighting factor. In any case, modifying  $E_b$  should lead to small improvements in the MAE of models since  $c_b$  is quite small.

With respect to the dependence of  $E_v$  on  $V_{r}$  our ideas for refining the model are motivated by the fact that the crossvalidated  $c_r$  value can be understood as the number of electrons

 $(N_e)$  donated by an O<sup>2-</sup> to its most reducible cation neighbors upon V<sub>O</sub> formation. Moreover, N<sub>e</sub> being approximately equal to 1.5 (see Figure 3a) and not 2 is related to how the cationreduction energy cost for creating  $V_{OS}$  gets split.  $V_r$  contributes about 75-80% of the cost, whereas  $\sim$ 15% of the cost is accounted for by  $E_{g}$  (which captures crystal field effects and any delocalization of electrons upon V<sub>O</sub> formation); this combination is why  $c_r + c_g$  adds up to 1.9. We hypothesize that the 10% remaining contribution comes from stability and bond-dissociation effects (where ions that do not undergo reduction can contribute). To explore further the role of electron delocalization in determining the  $c_r/c_g$  split across a wider range of the chemical phase space, one could calculate  $E_v$ for electronically diverse compounds whose anions neighbor O in the periodic table, such as metallic transition-metal phosphides for electrocatalysis,<sup>142–144</sup> semiconducting chalcogenides (sulfides and selenides for solar energy conver-sion),<sup>145-147</sup> and hybrid organic-inorganic metal halides (e.g., for optoelectronics).<sup>148–1</sup>

Finally, now we summarize some opportunities for extending the model by exploring other materials classes, including defect-defect interactions, and studying other defect types. First, we recommend testing (and, if necessary, retraining) the model on non-perovskite and non-ternary oxides. Second, for achievable Vo concentrations, defect-defect interactions may be non-negligible, thereby requiring the development of schemes to capture changes, for example, in the electrostatic repulsion between reduced and descreened cations and vacancy-vacancy interactions. Finally, we suggest that this approach be extended to other defect types such as bulk cation vacancies (which, e.g., affect the performance of LSCF cathodes for SOFCs<sup>151</sup>), charged vacancies and their transition levels (which, e.g., induce nonradiative recombination of photoexcited charge carriers in kesterite solar cells<sup>152</sup> and ABO<sub>3</sub> perovskites<sup>153</sup>), and surface vacancies (which, e.g., alter the electrical properties of  $SnO_2$  gas sensors<sup>154</sup>).

#### CONCLUSIONS

Metal oxide perovskites have received intense scrutiny for clean energy and computer memory applications due to their flexible structure, stability, wide composition space, and mixed ionic electronic conductivity. While exciting progress continues, a simple phenomenological explanation for the trends in oxygen vacancy  $(V_{\Omega})$  formation energies  $(E_{v})$ , which play a critical role in determining the performance of perovskite metal-oxide-based devices, has remained elusive. Here, we have introduced a machine-learned linear model that is based on  $E_{\rm v}$ values of  $ABO_3$  perovskites, where A = Ca, Sr, Ba, La, Ce and B = Ti, V, Cr, Mn, Fe, Co, Ni, in six lattice systems (monoclinic, orthorhombic, tetragonal, rhombohedral, hexagonal, and cubic). The model takes as inputs crystal bond dissociation energies, crystal reduction potentials, band gaps, and energies above the convex hull, which one can obtain from theoretical or experimental databases. Notably, we have devised crystal bond dissociation energies and crystal reduction potentials as the solid-state variants of their molecular and aqueous electrochemistry analogues, respectively, to systematically improve the description of bond dissociation and reduction of multivalent cations in solids. Additionally, we demonstrate that the model can be simplified, with acceptable losses in accuracy, such that only crystal bond dissociation energies and crystal reduction potentials are needed in scenarios where band gaps and energies above the convex hull are difficult to

estimate. Also, we predict BiFeO<sub>3</sub> and BiCoO<sub>3</sub> or a mixture might be viable candidates for solar thermochemical water and/or carbon dioxide splitting applications (intermediate  $E_{vv}$ , if the melting point can be pushed high enough: e.g., via Y-doping), with CeCrO<sub>3</sub> and ScCrO<sub>3</sub> as possible V<sub>O</sub>-resistant ferroelectrics (high  $E_v$ ), indicating the effect that our model can make in materials discovery. Adding further credibility to our model is our rediscovery of Co-based perovskites as robust solid oxide fuel cell cathodes (low  $E_v$ ). Finally, we present our perspectives on how to improve and extend the model, which already provides both accurate and efficient predictions for high-throughput screening and an intuitive and modular phenomenology for applications to metal oxide perovskites and beyond.

#### ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c05570.

Further details about the distribution of ABO<sub>3</sub> perovskite structures, distribution of neutral oxygen vacancy formation energies, convergence of oxygen vacancy formation energies in different crystal systems with respect to supercell size, distribution of energies above the convex hull, periodic trends in experimental cohesive energies, cell-size convergence of total energies for neutral atoms in their electronic ground states, theoretical and experimental cohesive energies across all metal oxides considered, correlation of experimental bond dissociation energies for neutral diatomic molecules and diatomic cations with respect to SCAN+Ucalculated crystal bond dissociation energies, specific trends in the SCAN+U-calculated and experimental crystal reduction potentials, and SCAN+U-calculated vs simplified-model-predicted oxygen vacancy formation energies (PDF)

Comma-separated-value files containing crystal structures, total energies, magnetic moments, charges, band gaps, and crystal features for all systems considered (ZIP)

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#### Notes

The authors declare no competing financial interest.

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