High-throughput determination of Hubbard $U$ and Hund $J$ values for transition metal oxides via the linear response formalism

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DFT+$U$ provides a convenient, cost-effective correction for the self-interaction error (SIE) that arises when describing correlated electronic states using conventional approximate density functional theory (DFT). The success of a DFT+$U(+J)$ calculation hinges on the accurate determination of its Hubbard $U$ and Hund’s $J$ parameters, and the linear response (LR) methodology has proven to be computationally effective and accurate for calculating these parameters. This study provides a high-throughput computational analysis of the $U$ and $J$ values for transition metal $d$-electron states in a representative set of over 2000 magnetic transition metal oxides (TMOs), providing a frame of reference for researchers who use DFT+$U$ to study transition metal oxides. In order to perform this high-throughput study, an atome workflow is developed for calculating $U$ and $J$ values automatically on massively parallel supercomputing architectures. To demonstrate an application of this workflow, the spin-canting magnetic structure and unit cell parameters of the multiferroic olivine LiNiPO$_4$ are calculated using the computed Hubbard $U$ and Hund $J$ values for Ni- and O- $p$ states, and are compared with experiment. Both the Ni-$d$ $U$ and $J$ corrections have a strong effect on the Ni-moment canting angle. Additionally, including a O-$p$ $U$ value results in a significantly improved agreement between the computed lattice parameters and experiment.

I. INTRODUCTION

Density functional theory (DFT) is a workhorse of computational materials science. However, the proper treatment of electronic exchange and correlation within the framework of DFT is a long-standing challenge [1]. Local density approximation (LDA) and generalized gradient (GGA) [2] functionals were developed to add exchange-correlation (XC) contributions to the energy functional within the Kohn–Sham (KS) formalism [3]. However, numerous studies have shown that these XC functionals have an associated self-interaction error (SIE) [1, 4, 5]. This shortcoming ultimately derives from the difficulty in quantifying exact exchange and correlation effects, without solving the many-body Schrödinger equation, using only density-based approximations.

Over the past couple of decades, DFT+$U$ has found favor as a method that strikes a reasonable balance between accuracy and computational cost, making it particularly suitable for high-throughput computation [6–10]. DFT+$U$ functionals add a correction to the conventional XC functional to account for the Coulombic interaction between localized electrons [4, 11]. In more recent studies, various researchers have explored extensions of DFT+$U$ with the goal of further correcting for static correlation effects and delocalization errors [12–14].

One drawback to DFT+$U$ type functionals is that one must first determine its associated parameters, the Hubbard $U$ and Hund $J$, and possibly also inter-site electronic interactions denoted as “$+IV$” [15–17]. The results of a DFT+$U$ calculation can quantitatively and even qualitatively change depending on these parameters, and so obtaining reliable values is of paramount importance.

This is as true for the Hund $J$ as it is for the Hubbard $U$, as we now explain. In this work, we primarily focus on the simplified rotationally invariant DFT+$U$ functional that has become very prominent since its introduction in Ref. 18. In this functional, the Hubbard $U$ and Hund $J$ are grouped in single effective Hubbard $U$ parameter, $U_{\text{eff}}$, defined as $U_{\text{eff}} = U - J$. This formalism assumes a spherical symmetry that results in the exclusion from the correction of the on-site exchange between opposite-spin electrons [10, 18, 19]. Notwithstanding, the reduction in the effective parameter by $J$ can be significant.

While the aforementioned approximation may seem more justifiable for systems with no magnetic order, in the case of magnetic systems it results in a lost opportunity to use the Hund $J$ to beneficially enhance the spin moments in simulated broken-symmetry ground states. Moreover, when we move to non-collinear magnetism, the spin texture of materials is particularly sensitive to screening interactions between spin channels [19–21]. In fact, magnetic exchange constants can be derived from...
the extended Hubbard model and estimated as ratios between $U$ and $J$ values [22]. The famous Hubbard model provides a simplified framework on which to explain the rich physics of correlated transition metal compounds [22]. Additionally, it has been shown that the Hund $J$ term is important for describing important physical phenomena, such as Jahn-Teller distortions [22, 23], emergent intra-atomic exchange, and the Kondo effect [24, 25]. Therefore, the introduction of explicit unlike-spin exchange corrections beyond simplified rotationally invariant DFT+$U$ is clearly of interest, and this requires the treatment of the Hund $J$ on the same footing as the Hubbard $U$.

A. Strategies for determining Hubbard parameters

A common approach for determining Hubbard $U$ values is to tune them such that some desired result — for example, the DFT+$U$ band gap, or a formation energy — matches its experimental value, or a value obtained via more accurate and computationally expensive beyond-DFT methods [26, 27]. There are several problems with this strategy. Firstly, it is not systematic: just because one result (e.g., the band gap) now matches experiment, this does not guarantee the same will be true for other observables (e.g., local magnetic moments). Indeed, there is a multitude of reasons why DFT may not match experiment, and it is wrong to rely on Hubbard corrections to correct for errors that do not arise from self-interaction [28]. Secondly, this strategy is not predictive: it relies on the existence of experimental/beyond-DFT data. This makes it particularly ill-suited to the prediction of novel materials and high-throughput studies.

Yet another difficulty that arises is the lack of transferability of Hubbard and Hund’s parameters. The conventional wisdom surrounding on-site corrections tends to reinforce the notion that localization equals correlation. Therefore, $+U$ corrections are applied to states determined by the orbital geometry (e.g., $d$ and $f$ orbitals). It is possible, using spectroscopy, to estimate Slater integrals over the Coulomb operator [18, 29] which, in turn, can be expressed in terms of $U$ and $J$ values [10, 30]. While this connection is physically motivated, localized states do not encompass all of the levels of correlation effects that are neglected by the specific DFT functional [13]. This perspective of $U$ and $J$ values as functional-specific, and not universal quantities, expands the definition of $U$ and $J$ from their initial inspiration from the Hubbard model, which treats $U$ and $J$ as intrinsic atomic properties. Indeed, it has been repeatedly shown that these parameters $U$ and $J$ are in fact very sensitive to the local chemical environment [31]. Even the specific pseudopotentials (PPs) [5] or the specific site occupation projection scheme [32] have a significant effect on the computed Hubbard $U$ values. The end result is that $U$ values (and by extension the albeit normally less environment-sensitive Hund’s $J$ values) are not transferable: they cannot be tabulated, and must always be determined on a case-by-case basis.

Having explored the numerous on-site corrections, and the drawbacks of fitting these parameters to experiment or beyond-DFT results, we will motivate the importance of computing Hubbard $U$ and Hund $J$ values within the DFT framework. Two primary methods for calculating Hubbard $U$ values in a self-contained fashion within DFT are the constrained random phase approximation (cRPA) [33, 34], and the linear response (LR) analysis of the constrained XC functional [4, 10]. In this study, we focus on the LR method due to its lower computational cost compared to existing cRPA methods, which are not yet appropriate for high-throughput applications. We also explore the effects on magnetic materials that exhibit a rich variety of noncollinear spin configurations, exemplified through the spin canting structure that was experimentally observed in olivine LiNiPO$_4$ [35].

The linear response method, as introduced for practical use by Cococcioni and coworkers [4], is founded on the idea that SIE can be related to the behaviour of the total energy as a function of the total occupation [36]. The energy ought to be piece-wise linear with respect to total site occupation numbers, but in fact for semi-local DFT XC functionals, the energy derivatives are erroneously continuous. Cococcioni and co-workers illustrated that the $+U$ correction can be interpreted as something that counteracts this erroneous curvature, locally for sub-spaces (where the interpretation becomes approximate). Crucially, the magnitude of the curvature can be directly measured from a DFT linear response calculation, allowing the value of $U$ to be determined accordingly. Unlike empirical fitting, this approach is (a) systematic, because the value of $U$ is derived directly as a measure of the underlying SIE present in the DFT calculation, and (b) it is predictive, because it only requires DFT calculations to extract the Hubbard parameters, and not experimental or beyond-DFT results.

B. Paper outline

The Materials Project is a web-based database that contains computed information on a vast range of materials, both known and predicted [37]. Among the various computational results it presents are Hubbard parameters $U_{\text{eff}}$. However, these current default $U_{\text{eff}}$ values were obtained by fitting DFT+$U$ energies to experimental formation energies for a selected number of redox reactions [31, 38]. This paper aims to replace these values with ones computed using linear response. In order to achieve this, we present a unified framework for computing on-site Hubbard and Hund’s corrections in a fully parallelized and automated computational workflow (which will be introduced in Section II). Using this workflow, we performed a high-throughput calculation of $U$ and $J$ values for a set of over two thousand transition-metal-containing compounds. This provides us with a
novel, big-picture point-of-reference for the sensitivity of $U$ and $J$ across a wide range of systems of varying chemistries and local chemical environments (Sections III A and III B). We then explore the effects of these Hubbard corrections on magnetic materials that exhibit a rich variety of noncollinear spin configurations, exemplified through the spin canting structure of olivine LiNiPO$_4$ (Section III C).

II. METHODS

A. The Hubbard functional

The Hubbard functional is a corrective functional, in the sense that it involves adding a corrective term $E_{\text{Hub}} - E_{\text{dc}}$ on top of some base functional $E_{\text{DFT}}$ (typically a local or semi-local functional), resulting in a total energy functional

$$
E_{\text{DFT}+U+J} = E_{\text{DFT}}[\rho] + E_{\text{Hub}} \left[ \left\{ n^\sigma_\gamma \right\} \right] - E_{\text{dc}} \left[ \left\{ n^\sigma_\gamma \right\} \right] = E_{\text{DFT}}[\rho] + E_{U/J} \left[ \left\{ n^\sigma_\gamma \right\} \right] \quad (1)
$$

The $(n^\sigma_\gamma)_{mm'} = \langle \varphi_{\gamma m} | \hat{\rho}^\sigma | \varphi_{\gamma m'} \rangle$ are matrices that represent the projection of the (spin-dependent) density operator onto Hubbard subspaces (indexed $\gamma$) defined by some set of orbitals $| \varphi_{\gamma m} \rangle$. These orbitals are typically atom-centred, fixed, spin-independent, localised, and orthonormal, often corresponding to the 3d or 4f subshell of a transition metal or lanthanide. The $n^\sigma_\gamma$ occupation numbers are the corresponding traces of $n^\sigma_\gamma$ matrices. The DFT+$U$ correction of Equation 1 adds a convex energy penalty to fractional occupations of these orbitals that in principle can counterbalance the SIE present in these Hubbard subspaces.

In the following paragraphs, we will provide a summary of some of the most well known formulations of DFT+$U$+$J$. We note that that because we are interested in the fully localized limit (FLL), we will not discuss extensions of DFT+$U$+$J$ to metallic systems, which employs an “around mean field” (AMF) methodology [10].

Starting from DFT+$U$+$J$ implementations of the highest complexity, and moving forward through increasing levels of simplification, we introduce the rotationally invariant implementation proposed by Liechtenstein et al. [30]. Within this flavor of DFT+$U$+$J$, $E_{\text{Hub}}$ and $E_{\text{dc}}$ take the following form

$$
E_{\text{Hub}} = \frac{1}{2} \sum_{\gamma, \sigma} U(m, m', m'', m''') (n^\gamma_{\sigma})_{m m'} (n^\gamma_{\sigma})_{m'' m'''} - \frac{1}{2} \sum_{\gamma, \sigma} U(m, m'', m', m') (n^\gamma_{\sigma})_{m m'} (n^\gamma_{\sigma})_{m'' m'} \quad (2)
$$

$$
E_{\text{dc}} = \sum_{\gamma} \frac{U^0}{2} n^\gamma (n^\gamma - 1) + \sum_{\gamma, \sigma} \frac{J^0}{2} n^\gamma (n^\gamma - 1) - \frac{1}{2} \sum_{\gamma, \sigma} J_{\gamma, \sigma} \left[ n^\gamma_{\sigma} (1 - n^\gamma_{\sigma}) \right] \quad (3)
$$

where $U$ contains the Coulomb integrals projected on the orbital basis, indicated by the associated $\{m\}$ quantum numbers [10, 12]. This correction is parameterized by both Hubbard $U^0$ and Hund $J_{\gamma}$ coupling constants through the double-counting energy contribution, $E_{\text{dc}}$.

Simplified versions of Equations 2 & 3 were proposed by Dudarev et al. [18], and later by Himmetoglu and coworkers [39], which approximate $U$ using Slater integrals, which can be parameterized through $U$ and $J$ values. There are many helpful explanations for this approximation, such as those summarized in Refs. 10 and 12.

In the spirit of following increasing levels of simplification, we will start with the Himmetoglu implementation [39], inspired by the work of Solovyev et al. [40]. Using the Slater integral parameterization of $U$ and $J$, it is possible to approximate and simplify $E_{U/J}$ from Equations 2 & 3 into the following

$$
E_{U/J} = E_{\text{Hub}} - E_{\text{dc}} = 
\sum_{\gamma, \sigma} \frac{U^0}{2} n^\gamma (1 - n^\gamma) + \sum_{\gamma, \sigma} \frac{J^0}{2} \text{Tr} \left[ n^\gamma_{\sigma} (1 - n^\gamma_{\sigma}) \right] \quad (4)
$$

A well known further simplification of Equation 4, notwithstanding that it substantially pre-dated the latter, is the formulation of DFT+$U$ put forth by Dudarev et al. [18] and given by

$$
E_U = E_{\text{Hub}} - E_{\text{dc}} = \sum_{\gamma, \sigma} \frac{U^0_{\text{eff}}}{2} \text{Tr} \left[ n^\gamma_{\sigma} (1 - n^\gamma_{\sigma}) \right] \quad (5)
$$

As discussed in the Introduction, this approximation arises by assuming spherical symmetry of the Coulomb interactions, $U$ [10, 12, 39]. Within the simplified Dudarev DFT+$U$, Equation 5, it has been demonstrated that the effective Hubbard U becomes $U^0_{\text{eff}} = U_{\gamma} - J_{\gamma}$ [10, 12, 18].

B. Hubbard $U$ and Hund’s $J$ spin-polarized linear response

In the linear-response approach, one measures the supposedly erroneous curvature in the total energy as a func-
tion of the subspace occupancy, and then chooses a value $U$ that counterbalances the observed curvature. Computing the energy curvature as a function of the subspace occupancy is usually impractical, so instead one transforms the curvature of the energy-versus-site occupancies $n_\gamma$ into a curvature with respect to the magnitude $v_\gamma$ of an on-site potential $\dot{v}_\gamma = \sum_{mm'} v_{\gamma m'} |\varphi_{\gamma m'}\rangle \langle \varphi_{\gamma m'}|$. The energy functional is then given by

$$E[\{v_\gamma\}] = \min_{\rho(\mathbf{r})} \left\{ E[\rho(\mathbf{r})] + \sum_\gamma v_\gamma n_\gamma \right\}$$

where $v_\gamma$ is the on-site potential at site $\gamma$. From this energy functional, one computes the response matrices

$$\chi_{\gamma \gamma'} = \frac{\partial n_\gamma}{\partial v_{\gamma'}}.$$

Thus far we have used a general index “$\gamma$” to represent each site. Conventionally, this index refers purely to the atom $\gamma$ on which the Hubbard site is centered. In this case, the Hubbard parameter for that subspace is given by

$$U_\gamma = \left( \chi_0^{-1} - \chi^{-1} \right)_{\gamma \gamma}$$

where $\chi$ and $\chi_0$ are the interacting, (or self-consistent) and non-interacting (or non-self consistent) response matrices [4, 10].

The above strategy does not delineate between spin channels: during the linear-response calculations the spin-up and spin-down channels are perturbed simultaneously by the same amount, i.e., $v_{\gamma}^\uparrow = v_{\gamma}^\downarrow$ and we only observe the change in total occupancy $n_\gamma = n_{\gamma}^\uparrow + n_{\gamma}^\downarrow$. If we want to calculate $J$, one must instead consider the spin-dependent perturbation

$$\dot{v}_\gamma^\sigma = \left\{ + \sum_{mm'} v_{\gamma m'} \varphi_{\gamma m'} \langle \varphi_{\gamma m'} | \right|_{\sigma = \uparrow} - \sum_{mm'} v_{\gamma m'} \varphi_{\gamma m'} \langle \varphi_{\gamma m'} | \right|_{\sigma = \downarrow}$$

and then construct a second set of response matrices which then relate to $J$ in a completely parallel approach to the calculation of $U$ in 8.

A separate but ultimately equivalent strategy is to treat the spin channels separately [5, 41]. In this case a general index runs over both the atom index $\gamma = \{1, \ldots, N\}$ and also the two spin channels $\sigma = \{\uparrow, \downarrow\}$. In this case the response matrices of Equation 7 become rank-four tensors, i.e.,

$$\chi^{\sigma \sigma'}_{\gamma \gamma'} = \frac{\partial n^\sigma_{\gamma}}{\partial v^\sigma_{\gamma'}}.$$

and now the equivalent of Equation 8 is

$$f_{\gamma \gamma'}^{\sigma \sigma'} = \left( \chi_0^{-1} - \chi^{-1} \right)_{\gamma \gamma'}^{\sigma \sigma'}$$

C. Implementation of linear response within a high-throughput workflow

The linear response method was implemented as a workflow within the high-throughput atomate framework [42]. The workflow allows the user to compute Hubbard $U$ and Hund $J$ values using either a spin-polarized or a non-spin-polarized response. In addition to screening between spin channels, the implementation provides the straightforward extension to multiple levels of screening, including inter-site and inter-spin-channel responses [5]. A more detailed explanation of how these screening matrices are computed is provided in Appendix A.

All of the individual calculations within this workflow were performed with VASP (Vienna ab initio Simulation Package) [43], a plane-wave DFT code. The PBE exchange-correlation functional was used throughout as the base functional [44]. Unless otherwise stated we use PAW PBE pseudopotentials (PPs), which are the default PPs for the pymatgen input sets for VASP [45]. In this regard, our work supplements the high-throughput work of Bennett et al. [46] where ultrasoft pseudopotentials (USPPs) were used to reduce computational cost in high-throughput computations [46], mirroring early foundational studies on the linear response method [4, 15].

We have used an automatic k-point generation scheme that uses 50 k-points per reciprocal angstrom, and a cutoff energy of 520 eV. The full set of input parameters can be found in the HubbardHundLinRespSet in the atomate repository [47], and the derived VASP input sets in the pymatgen repository [45]. For the linear response analysis, the on-site applied potential $v_{1\sigma}$ range was from $-0.2$ eV to $+0.2$ eV ($-0.05$ eV to $+0.05$ eV for the periodic table data set) sampled at nine points at uniform intervals.
TABLE I: Comparison of computed $U_{\text{eff}}$ in the present work with values used by the Materials Project [31, 38].

| element | mean $U_{\text{eff}}$ (eV) | $U_{\text{eff}}^{\text{MP}}$ (eV) | diff. (eV) |
|---------|-----------------------------|-------------------------------|-----------|
| Co      | 4.430 ± 1.474               | 3.32                          | 1.110     |
| Cr      | 2.425 ± 0.472               | 3.7                           | -1.275    |
| Fe      | 4.108 ± 1.322               | 5.3                           | -1.192    |
| Mn      | 4.135 ± 0.724               | 3.9                           | 0.235     |
| Mo      | 1.911 ± 0.318               | 4.38                          | -2.469    |
| Ni      | 5.258 ± 0.773               | 6.2                           | -0.942    |
| V       | 3.060 ± 0.673               | 3.25                          | -0.190    |
| W       | 1.461 ± 0.218               | 6.2                           | -4.739    |

III. RESULTS

Hubbard $U$ and Hund $J$ values were calculated for over two thousand transition metal oxides using the linear response workflow implemented in atomate. The majority of the calculations corresponded to materials containing Mn-$d$, Fe-$d$, and/or Ni-$d$ species. All the systems studied were previously predicted by Ref. 48 to have a collinear magnetic ground-state using a separate high-throughput workflow. That work used the empirical Hubbard $U$ values reported on the Materials Project.

In addition, a representative set of O-$p$ responses were calculated and analyzed. It is less common to include Hubbard corrections to oxygen 2$p$ states. However, an appreciable number of studies have shown how O-$p$ on-site corrections have improved the agreement with experimentally measured bond lengths between oxygen and transition metal species [5, 49–52]. It is perhaps less intuitive to apply spin-polarized Hund $J$ parameters to oxygen sites, because O-$p$ states are conventionally not included in effective models for magnetism. However, while oxygen atoms do not develop magnetic moments, early studies have demonstrated theoretically and computationally that O-$p$ states mediate the antiferromagnetic superexchange interaction in transition metal oxides, such as MnO [22, 53, 54].

A. Periodic table sample set

Figure 1 displays two periodic tables containing the distributions of computed Hubbard $U$ and Hund’s $J$ values for each transition metal element (and oxygen) computed for different structures within the database. In Table I, values obtained in this study are listed alongside the standard $U$ values employed by the Materials Project [31, 38]. Those values were determined using the procedure outlined by Wang et al. [56] which finds a $U$ value that minimizes the error in formation energy for several representative redox couples. Due to the limited amount of experimental data available, these $U$ values are determined with only experimental data from a single redox couple (Co, Cr, Mo, Ni, and W) or two redox couples (Fe, Mn, and V). Therefore, it is possible or likely that these $U$ values are not appropriate for a more general system containing these elements. Nevertheless, the MP $U$ values are found to be the same as the $U$ values in the present work within the standard deviation for most elements (Co, Fe, Mn, and V) or slightly outside the value in the present work (Ni). Exceptions are Cr, Mo, and W, with the largest, notable discrepancy of 4.739 eV for W.

To evaluate the impact of these discrepancies, compounds containing W from a dataset of experimental formation energies [57] used by the Materials Project were taken and relaxed using the new $U_{\text{eff}}$ value for W from the present work but with all other calculation settings kept consistent with standard Materials Project settings, to obtain a new set of computed energies. These energies substantially lowered the correction introduced in Ref. 57 for W from -4.437 eV/atom to 0.12 eV/atom, suggesting that the newer $U_{\text{eff}}$ is indeed more appropriate for the calculation of formation energies.

We stress that these values are not transferable to other studies, which use DFT implementations in other codes. QUANTUM ESPRESSO and ABINIT use localized projections that are different from the projector augmented wave (PAW) method implemented in VASP [32].

B. Focused study on Mn-$d$, Fe-$d$, Ni-$d$, and O-$p$, including the reason for large O-$p$ Hubbard $U$ values

We now present a more detailed study on materials containing Mn-$d$, Fe-$d$, Ni-$d$, and O-$p$ Hubbard sites. For these systems, the distributions of the computed Hubbard $U$ and Hund $J$ values are provided in Figure 2. The variations in $U$ and $J$ values calculated for these three species is immediately apparent, with a range on the order of approximately 1 to 2 eV. These distributions reflect the intrinsic screening environment dependence of the calculated value for a given element. At this point, we note only their apparently universal unimodality (single peak) and the near-general decrease in $U$ with chemical period within a given group, however we will return presently to a more physically and chemically motivated observation. In Table II we list for comparison the $U$ values currently used in Materials project (fitted empirically) as well as a range of $U$ values found for a set of spinels and olivines by Zhou and co-workers (calculated via self-consistent linear response) [31].

We find that O-$p$ exhibits the largest associated Hubbard $U$ value of approximately 10 eV, which agrees with the linear response results from a previous study using a different code and somewhat different linear-response formalism [5]. While large oxygen Hubbard $U$ values may seem surprising within a strongly correlated materials context, it has become more accepted in recent years within first-principles solid-state chemistry that oxygen 2$p$ orbitals can warrant, both by direct calculation and by necessity (when resorting to fitting), a remarkably high $U$ value in DFT+$U$. 


FIG. 1: Periodic tables of Hubbard $U$ and Hund $J$ values computed for a representative set of transition metal oxides. The color map indicates the mean value computed for each element over each material. The materials used in the creation of these periodic tables were selectively chosen: noting that many databases, including the ICSD, contain a growing number of hypothetical materials which may or may not be realizable, we selected materials that are well-studied and exhibit more than two ICSD IDs each. Furthermore, to remove cross-correlations between magnetic elements, we also require that these compounds only contain a single $d$-block element (occupying a single symmetrically-equivalent site) with no $f$-block species. Ultimately this data corresponds to the $U$ and $J$ values for over 800 materials, and are distributed over the transition metal species. A more detailed table containing data on the distribution of values is included in Appendix C. The plotted distributions of $U/J$ values are generated using a Gaussian kernel-density estimator implemented in scipy [55].
We will now attempt to motivate and explain this phenomenon. We note from the outset that the element projector orbital profile plays a complicating role in the following analysis. In general, we observe that the diagonal elements of the $\chi_0$ non-interacting response matrix are of roughly the same magnitude for both TM-$d$ and O-$p$ sites. The non-self-consistent response can be interpreted as the response due to non-interacting response effects at a site due to its surroundings [10], and thus it can be understood as a property primarily of the environment of the atom under scrutiny. Then, unless screening is very short ranged as it may be in a very wide-gap insulator, this quantity may be said to be somewhat similar, on average, for metal and oxygen ions in an oxide. Thereby, the chemical trends in the Hubbard $U$ arise mostly in the interacting response.

Next, we note that the O-$p$ interacting response $\chi$ tends to be less than half of that of the interacting TM-$d$ response. This indicates that $-\chi^{-1} = d^2E/dn_i^2$, the curvature of the total energy versus occupation, $n_i$, is greater for O-$p$ states. This greater curvature versus occupation can be explained, we propose, in terms of known trends in the chemical hardness, i.e., the second chemical potential, i.e., the derivative of the chemical potential with respect to total charge at fixed external potential. We note, in passing, that some authors choose to define the chemical hardness as half of that for historical reasons, but we suppress that here. Specifically, we can focus on the discretized (three-point) approximation to the global chemical hardness [58], namely

$$
\nu \equiv d^2E/dN^2 \approx E(N-1) - 2E(N) + E(N+1)
$$

$$
= [E(N+1) - E(N)] - [E(N) - E(N-1)]
$$

$$
\equiv E_i - E_a \equiv Eg,
$$

(12)

which is nothing but the fundamental band-gap. This is a quantity that has been tabulated many times, and using the results of Ref. 59 we find that for atomic oxygen its value is 11.2 eV, compared to that of the transition metal atoms, where it ranges from 5.8 eV (Ti & Zr) to 8.0 eV (Mn) if we exclude the often problematic zinc group, where it reaches 11.6 eV. This mirrors and explains the observed relatively large first-principles Hubbard $U$ value for oxygen 2p states predicted in this and several previous studies.

Ultimately, we conclude that the Hubbard $U$ may be interpreted as the subspace-projected, environment screened chemical hardness, and more precisely as only the interaction (e.g., Hartree, exchange, correlation, and perhaps other terms like implicit solvent and PAW potential) part of that. It is in the interaction part that most of the chemical trends appear to arise in practice. For subspaces projecting heavily at both band-edges, as in normal DFT+$U$ practice, the $U$ clearly inherits chemical trends from the chemical hardness (fundamental gap) of the atom that it resides upon. This is higher for a higher atomic ionization energy $E_i$ (that of oxygen is generally around twice that of transition metals) and higher also for a more negative electron affinity $E_a$ (that of oxygen is more negative than that of most but not all transition metals). By and large, both quantities are well known to increase in magnitude as we move ‘up and right’ in the periodic table, and this same broad trend is reflected in our periodic table of Hubbard $U$ values.

When a DFT+$U$ subspace projects only onto one or other band edge, as seems more commonly the case for charge-transfer insulators, then then the trend in only one of the ionization energy and electron affinity will be very relevant to the trends in $U$. In the case of oxygen 2p orbitals projectors, due to the electronegativity of oxygen typically there will be little weight at the conduction band edge, and so it is the (particularly clear) trend in ionization energy that drives the relatively large $U$ value for oxygen. Indeed, if this argument holds then one would guess that the oxygen 2p $U$ value is roughly twice that of an average transition-metal d subspace, which turns out to be the case from first principles linear response.

The Hund’s $J$, within the present formalism, may be interpreted as an analogue for the spin degree of freedom, and specifically as minus (by a convention thought to originate with Ising) the interaction part of the subspace-projected, environment screened spin-hardness, even the global atomic version of which [60] has been a much less thoroughly studied quantity. The effect of choosing whether these subspace charge (spin) hardness quantities, the $U$ and $J$, are calculated in a fully relaxed manner, or with with a simulated fixed spin (charge), is explored in our comparison between conventional (scaled) and constrained (simple) spin-polarized linear response, respectively, below.

| Species | mean computed $U$ (eV) | mean computed $J$ (eV) | mean computed $U_{\text{eff}} = U - J$ (eV) | reported $U_{\text{eff}}$ MP [38] (eV) | reported $U_{\text{eff}}$ range [31] (eV) |
|---------|------------------------|------------------------|---------------------------------------------|--------------------------------------|--------------------------------------|
| Mn-$d$  | 4.953 ± 0.635          | 0.520 ± 0.156          | 4.433 ± 0.654                               | 3.9                                 | 3.60 – 5.09                          |
| Fe-$d$  | 4.936 ± 0.700          | 0.177 ± 0.367          | 4.759 ± 0.790                               | 5.3                                 | 3.71 – 4.90                          |
| Ni-$d$  | 5.622 ± 1.221          | 0.399 ± 0.434          | 5.223 ± 1.296                               | 6.2                                 | 5.10 – 6.93                          |
| O-$p$   | 10.241 ± 0.910         | 1.447 ± 0.171          | 8.794 ± 0.926                               | N/A                                 | N/A                                  |
FIG. 2: Distributions of Hubbard $U$ and Hund $J$ values computed using the linear response method; For the sub-figures (a), (b), and (c) that correspond to $d$-electron TM site corrections, the $U$ and $J$ values are plotted against the DFT (no $+U$+$J$ correction) computed site magnetic $m_l$, where $m_l$ and $m_d$ are equivalent because $d$ occupations have a corresponding $l = 2$ angular momentum quantum number. The $O$-$p$ $U$ and $J$ values in sub-figure (d) are plotted against $n_l$ ($n_1$ or $n$-$p$) total site occupations. The number of samples for on-site correction values for Mn-$d$, Fe-$d$, Ni-$d$, and O-$p$ are 285, 248, 149, and 206, respectively.

In order to explore trends in the distribution of $U$ and $J$ values, we have plotted these on-site corrections in scatter plots within Figure 2. These plots illustrate the relationship between $U$ and $J$ values with respect to site occupations. For transition metal species, we plot $U$ and $J$ versus $m_{l=2}$, the “$d$” component of the projected moment, $m$, denoted as “$m_{l=2}$.” These moment values are those output by VASP as the difference of up and down spin site occupancy numbers computed using PAW site-projection operators. Because the oxygen atoms do not have an associated magnetic moment, we plot O-$p$ Hubbard $U$ and Hund $J$ versus $n_{l=1}$ occupations on oxygen sites.

We should stress that the values of “$m_l$” and “$n_l$” are only computed from the calculation without the $+U$ correction. One reason for using the bare PBE computed $m_l$ and $n_l$ is that these occupations should be independent from the applied Hubbard $U$ or Hund $J$ values. This would offer the “bare” $m$, as well as $n$, as a possible predictors of $U$ and $J$ values. However, it is important to note that these occupations could change significantly with applied $U$ and $J$ values [5, 61, 62].

There is an apparent clustering of data points at different on-site $m_l$ magnetizations in Figures 2a, 2b, and
This grouping at different on-site magnetization values is most likely due to different spin and charge states dependent on the underlying chemistry. We also observe a larger range of \( U \) and \( J \) values for higher values of \( m_{\uparrow} \), which is due to the coupling between highly spin-polarized states to on-site Coulomb screening for TM species. As would be expected, we see similar trends for \( J \), a measure of the screened interaction between spin channels.

For the Mn-\( d \) and Ni-\( d \) distributions in Figures 2a and 2c, a stark clustering of data-points is evident at particular intervals of \( m_{\downarrow}=2 \). In both cases, the clusters that lie at the associated maximum computed \( m_{\uparrow} \) fall off and exhibit a negative slope trend with the magnitude of the site moment. This is likely due to the fact that \( m_{\uparrow} \) is highly dependent on the local chemical environment, which will govern the energy curvature over spin occupations, which is directly related to \( U \) and \( J \) within linear response [4]. The clear trend for the manganese may be due to the strong tri-modal distribution of Mn magnetic moments seen in Figure 1 of Ref. 48. The “stable” magnetic configurations from this study were used in the LR analysis, therefore a similar statistical distribution should hold for the subset of structures used in this LR analysis.

The trends of the data points for Hubbard \( U \) and Hund \( J \) values in Figure 2d appear to show a downward trend for \( U \) versus \( p \)-occupation numbers, \( n_{\downarrow}=1 \), and a slower, upward trend for \( J \) versus \( n_{\downarrow}=1 \). We expect that the \( n_{\downarrow}=1 \) occupations will be strongly dependent on the oxidation/reduction state of oxygen atoms. Due to the nature of TM-O bonding in these oxides, and their generally greater electronegativity, the oxygen atoms will tend to maximize their valence. Therefore, building on the previous explanation of the magnitude of O-\( p \) \( U \) values based on chemical hardness and specifically the more relevant ionization potential component of that, the higher electron count for oxygen corresponds to a lower ionization potential, and therefore to a reduced Hubbard \( U \), as observed.

In an attempt to more robustly tease apart these observed trends, we performed a rudimentary random forest regression test on the data set, ultimately in an attempt to predict the on-site corrections \( U \) and \( J \) from the input crystal structures and site properties. We used the random forest regression algorithm as implemented in scikit-learn. The input quantities supplied to the random forest regressor consisted of the corresponding PBE-computed \( m_{\uparrow} \) and \( n_{\downarrow} \) - without on-site corrections, as well as the oxidation state estimated using the bond valence method [63], and finally a selection of relevant site features as formulated by \[64, 65\]. The OFM encodes the orbital character of the surrounding chemical environment. For more information on this method please refer to Ref. 65. The OFM functionality is not implemented in matminer or pymatgen. We were motivated to test the vectorized OFM by the chemical intuition that on-site Hubbard \( U \) and Hund \( J \) values are very sensitive to the local chemical environment. Additionally, the OFM has demonstrated success in predicting DFT-computed magnetic moments in the past [65]. Furthermore, the OFM nearest-neighbor contributions are weighted according to the geometry of the Voronoi cell, which could possibly provide information beyond the relative importance of the Voronoi matminer featurizer. However, the on-site magnetization for Mn, Fe, and Ni, respectively, had an importance of at least ten percent more than any of the other local chemical environment descriptors.

The correlation between on-site corrections and projected site moments is not surprising. After all, previous studies have explored the connection between charge states of transition metal species and the integrated net spin calculated from DFT [67–69]. The integrated atomic spin moment can be directly linked to the charge state of transition metal species via magnetochemistry rules. In fact, recent studies show that the magnetic moment is often the most convenient and reliable indicator of charge states [67].

### 1. Conventional vs. constrained linear response

In introducing the linear response theory in Section II B, we mentioned that there are two possible schemes for computing \( U \) and \( J \): “conventional” and “constrained” linear response, where in the latter case the linear response is performed in such a way that the magnetic moment (occupation) is held fixed while measuring the curvature with respect to the occupation (magnetic moment). While arguments can be made as to theoretically which approach is the most valid (a topic which is the subject of ongoing research), this dataset presents an opportunity to evaluate how much this choice will practically affect the resulting Hubbard and Hund’s parameters.

For the majority of the computed \( U \) and \( J \) values using
these two methods, the difference between the two strategies fell within their computed uncertainty. However, we observed a significant deviation from $y = x$ behavior for the computed $U$ values for iron Hubbard $U$ values shown in Figure 3. The width of this distribution is greater than 1 eV for $U$ in some regions, which is enough to affect computed physical properties \[4, 61\].

2. Dependence on structure and magnetic state

For some input magnetic structures, the magnetic configuration changed while applying the on-site potentials during the linear response analysis. Our hypothesis is that the input magnetic structure corresponds to a local minimum configuration, or possibly a metastable state. Therefore, in our analysis, we screen out these structures with the intent that these systems will be studied in the future using a self-consistent approach to calculating on-site corrections.

In order to test the sensitivity of $U$ and $J$ values to the input structure, we perform a self-consistent linear response study of antiferromagnetic NiO, which is provided in the Supplementary Information. Each iteration consists geometry optimization of cell shape, followed by a linear response calculation of $U$ and $J$ values. These on-site correction values are then used in the next subsequent geometry optimization step. Self-consistency is achieved once the $U$ and $J$ values fall within their corresponding uncertainty values. Starting from the input structure — which was optimized using the current default Materials Project $U$ values \[38\] — convergence was achieved after only two iterations.

It has been well established in previous studies that $U$ values should be computed self-consistently with geometry optimization \[61\]. As demonstrated from the experiments with antiferromagnetic NiO in the Supplementary Information section, the Hund $J$ values should be calculated self-consistently, in addition to Hubbard $U$ values. In this self-consistency study, $J$ had the largest associated change over convergence relative to the value itself. Due to the coupling between Hund $J$ and magnetic exchange \[22\], it is possible that both magnetic and structural features should be included in the self-consistency cycle. Within the atomate framework, it would be possible to incorporate a workflow that wraps the workflow developed in this study, in order to alternate linear response calculations with geometry relaxation until self-consistency is achieved.

2. Dependence on structure and magnetic state

C. Case study: LiNiPO$_4$

We now present a detailed study on the olivine LiNiPO$_4$, designed to test the results produced by the linear response workflow. Previous GGA+$U$ and GGA+$U$+$J$ studies have attempted to reproduce the experimentally-observed spin-canting structure and unit cell shape as shown in Figure 4 \[19, 31, 35\].

We calculated $U$ and $J$ for this system via spin-polarized linear response. The spin-polarized linear response method introduced in Section II B can be generalized to noncollinear DFT using the relationship between spin-density occupations and the magnitude of the magnetic moment: $n_\uparrow = \frac{1}{2} (n + |\vec{m}|)$ and $n_\downarrow = \frac{1}{2} (n - |\vec{m}|)$ \[21\]. For comparison, we also performed a collinear calculation, where the magnetic configuration for LiNiPO$_4$ was obtained by projecting the canted noncollinear structure shown in Figure 4 along the $z$-direction. In addition to one unit cell of the the collinear antiferromangetic (AFM) configuration, a linear response analysis was performed.
TABLE III: Hubbard and Hund results for Ni-d in LiNiPO$_4$ (Atom-wise screening)

| cell          | magnetism | $U$ (eV) | $J$ (eV) |
|--------------|-----------|----------|----------|
| $1 \times 1 \times 1$ | collinear | 5.43 ± 0.16 | 0.38 ± 0.07 |
| $1 \times 2 \times 2$ | collinear | 5.44 ± 0.24 | 0.54 ± 0.07 |
| $1 \times 1 \times 1$ | non-collinear | 5.09 ± 0.15 | 0.42 ± 0.05 |

on a $1 \times 2 \times 2$ supercell. Table III summarizes the results of the computed Hubbard $U$ and Hund $J$ values. From this table, it is evident that the $U$ value is significantly smaller in magnitude with the inclusion of spin-orbit coupling. A possible justification for this behavior is the introduction of orbital contributions to the total localized magnetic moments with the inclusion of spin-orbit coupling [72, 71].

1. Canting angle exploration

In order to explore the canting angle experimentally observed for LiNiPO$_4$ [35], we calculated the energy as a function of constrained canting angle. The noncollinear magnetic constraints were performed in VASP in accordance with the method developed by Ma and Dudarev [72]. We used the experimentally derived spin canting structure as a reference provided by the Bilbao Crystallographic Server, as shown in Figure 4 [35, 70]. The energy versus canting angle curve is shown in Figure 5a. We found that the stable canting direction is in the opposite direction to the experimentally measured canting angle. However, this discrepancy with experiment was limited to the canting direction; the computed stable magnetic structure still obeyed the symmetry of the Pm$\bar{3}$a magnetic space group.

Similarly to the work by Bousquet and Spaldin [19], we observe an increasing canting angle with Hund $J$ value. Interestingly, adding a $U$ and $J$ correction to O-p results in a slightly decreased stable canting angle. However, we find that in all cases, the computed stable canting angle is significantly less than the experimentally measured canting angle of 7.8 degrees [35].

The constraining effective site magnetic field, $\vec{H}^\text{eff}$, can be described as the following

$$
\vec{H}^\text{eff}_i = 2\lambda \left[ \vec{M}_i - \hat{M}_i \left( \hat{M}_i \cdot \vec{M}_i \right) \right]
$$

where $\vec{M}_i$ are the integrated magnetic moments at site $i$, and $\hat{M}_i$ are the unit vectors pointing in the individual site constraining directions [72]. The $z$ component of the constraining field (in the direction of canting), $H^\text{eff}_{1,z}$, is plotted versus the constraining angle in Figure 5b. We see that where $H^\text{eff}_{1,z}$ changes sign corresponds to the minimum of Figure 5a.

2. Effect of $U$ and $J$ values on geometry optimization

While the addition of Hubbard and Hund parameters go some way to addressing the canting angle of LiNiO$_4$, introducing these terms can also alter the geometry of the system. To explore this effect, we performed structural relaxations of the system with various combinations of Hubbard and Hund’s corrections. In each of these structural relaxation calculations, a maximum force tolerance was used of 10 meV/Å. All runs included spin-orbit coupling, and were constrained to the 7.8 degrees experimentally observed canting angle.

Table IV lists the optimized unit cell parameters and volume, compared with the experimentally measured geometry [35]. For both the PBE+$U_{\text{eff}}$ and PBE+$U$+$J$ schemes, adding corrections to the Ni-d space worsens the geometry relative to the uncorrected PBE geometry (as earlier observed by Zhou and co-workers [31]). However, the further addition of corrections to the O-p subspace reduces the errors by three-fold, resulting in geometries that are closest to experiment. This is similar to observations in other studies when applying corrections to O-p subspaces [5]. We note that applying a $+J$ correction to non-magnetic O-p states is unconventional. However, it should be stressed that the projected magnetic moments on LiNiPO$_4$ remain just below 0.01 $\mu_B$, with and without on-site corrections to O-2p states. Meanwhile, we can see that adding a $+J$ parameter does not significantly alter the cell parameters.

The Hubbard $U$ and Hund $J$ values used in this study of LiNiPO$_4$ include those calculated using linear response, which are approximations of the values that are reported in Table III. Additionally, we tested the Ni-d $U$/$J$ values used in Ref. 19, in order to compare with previous computational studies of the magnetic structure of LiNiPO$_4$.

3. Discussion on TM-O bond length versus $U$, $J$, and $V$ corrections

Table IV also presents the change in mean Ni-O bond length between nearest-neighbor pairs for various on-site corrections. For the Ni-O bond length it is the same story as for the cell parameters: applying $U$ and $J$ to the Ni-d sites worsens the results relative to the PBE result, but by applying corrections to the O-p channels we obtain bond lengths that are in closer agreement with experiment. In Ref. 5, some of us attempted to rationalize this trend in the computed bond length between transition metal species and oxygen anions and how it improves with the introduction of corrections to the O-p subspace [5]. We suggested that when $+U$ is added to the Ni-d subspace the resulting potential shift disrupts hybridization between the Ni-d and O-p orbitals, weakening the bonding between these two elements (and thus leading to bond lengthening). By applying corrections to the O-p re-aligns these two subspaces and allows them to
“re-hybridize”.

In an attempt to more thoroughly explore this reasoning, Figure 6 provides a comparison for the projected density of states (DOS) of LiNiPO$_4$ for PBE and PBE+$U+J$ (with and without corrections to O-$p$). It is difficult to discern re-hybridization from DOS plots alone.

Without an explicit quantification of hybridization effects, we have added a derivation in the Supplementary Information that presents a mathematical expression of the forces acting on ions due to $+U+V$ corrections. This result is an extension of the theory put forth by Matteo Cococcioni in Chapter 4, Section 4.1 of Ref. 10. We argue that in quantifying the forces on TM-O bond lengths due to on-site corrections, it is possible to show that the force contributions due to both $+U$ and $+V_{\gamma\gamma'}$ can, and should, be treated on the same footing, where $\gamma$ and $\gamma'$ correspond to atomic sites. It isn’t possible to definitively state the comparative magnitude, or sign, of these force contributions without additional calculations or simplifications based on physical intuition. However, the result suggests that the forces on TM-O bond lengths due to O-$p$ $U$ values will have a comparative magnitude to the forces due to inter-site Coulomb corrections due to $+V$.

In the Supplementary Information, we further hypothesize the sign of these force contributions, starting from a DFT geometry-optimized structure without on-site corrections. Using these assumptions, which are based on computational trends in bulk TMOs, we conclude that both applying a $+U$ correction to the O-$p$ manifold and a $+V$ between TM and O states combine to mitigate the overestimation of TM-O bond length that arises when only applying $+U$ to localized states around the TM species.

IV. CONCLUSIONS

This study provides a high-throughput atomate framework for calculating Hubbard $U$ and Hund’s $J$ values. Using the spin-polarized linear-response methodology [5], we generated a database of $U$ and $J$ values for over two thousand transition-metal-containing materials. This enabled the creation of a “periodic table” of $U$ and $J$ values, where for each element we observe a distribution of Hubbard $U$ and Hund’s $J$ values. These distributions exhibit...
FIG. 6: Projected electronic density of states for LiNiPO$_4$ (calculated using experimental unit cell [35]) without Hubbard or Hund corrections applied, as well as $+U$ and $+J$ applied to Ni-d channels, and both Ni-d and O-p states, respectively.

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Credit Taxonomy

We highlight the author contributions to this study using the CRediT taxonomy. Guy C. Moore: Conceptualization, Methodology, Software, Validation, Formal analysis, Investigation, Data Curation, Writing - Original Draft, Writing - Review & Editing, Visualization Matthew K. Horton: Conceptualization, Software, Validation, Investigation, Writing - Review & Edit-
Appendix A: Screening matrix inversions

Below are the matrix representations of the response matrices at each level of screening outlined by Linscott and others for a system with two Hubbard sites [5].

Point-wise $1 \times 1$ screening:

$$\chi^{-1} = \begin{pmatrix} 1/\chi_{11} & 0 \\ 0 & 1/\chi_{22} \end{pmatrix}$$

(A1)

Atom-wise (conventional) $2 \times 2$ screening:

$$\chi^{-1} = \begin{pmatrix} \chi_{11} & \chi_{12} \\ \chi_{21} & \chi_{22} \end{pmatrix}^{-1}$$

(A2)

We can extend this formalism to the multiple site (multiple-site) responses by considering the response matrix for two sites, $\chi_{ij}$, where $i$ and $j$ are the site indices.

Point-wise screening:

$$\chi^{-1} = \begin{pmatrix} 1/\chi_{1111} & 0 & 0 & 0 \\ 0 & 1/\chi_{1112} & 0 & 0 \\ 0 & 0 & 1/\chi_{1122} & 0 \\ 0 & 0 & 0 & 1/\chi_{1212} \end{pmatrix}$$

(A3)

Atom-wise (conventional) screening:

$$\chi^{-1} = \begin{pmatrix} \chi_{1111}^{-1} & 0 & 0 & 0 \\ 0 & \chi_{1112}^{-1} & 0 & 0 \\ 0 & 0 & \chi_{1122}^{-1} & 0 \\ 0 & 0 & 0 & \chi_{1212}^{-1} \end{pmatrix}$$

(A4)

Full screening:

$$\chi^{-1} = \begin{pmatrix} \chi_{1111}^{-1} & \chi_{1112}^{-1} & \chi_{1122}^{-1} & \chi_{1212}^{-1} \\ \chi_{1112}^{-1} & \chi_{1122}^{-1} & \chi_{1212}^{-1} & \chi_{1222}^{-1} \\ \chi_{1122}^{-1} & \chi_{1212}^{-1} & \chi_{1222}^{-1} & \chi_{2222}^{-1} \\ \chi_{1212}^{-1} & \chi_{1222}^{-1} & \chi_{2222}^{-1} & \chi_{2222}^{-1} \end{pmatrix}$$

(A5)

We note that it is important when performing a linear response calculation to construct a $2N \times 2N$ response matrix where $N$ is the number of Hubbard sites (or $N \times N$ in the case of non-spin-polarized linear response). For bulk systems often several Hubbard sites will be equivalent, and one can save computational time by performing linear response calculations for the set of inequivalent sites, and then populating the response matrix for all equivalent Hubbard-site pairs.

Appendix B: Post-processing & uncertainty quantification

In order to extract the response matrices from the raw DFT data, curve fitting was performed using a least-squares polynomial fit implemented in numpy [73]. The uncertainty associated with each computed slope was obtained from the covariance matrix produced as a result of the least-squares fit. These uncertainty values were then utilized to determine the errors associated with the Hubbard $U$ and Hund $J$ values. The error quantification was performed by computing the propagation of uncertainty based on the Jacobian of each scaling formula for Hubbard $U$ and Hund $J$. This method for error propagation is general to multiple levels of screening between spin, site, and orbital responses.

We begin by considering the following screening matrix introduced in Equation 11, from which Hubbard $U$ and Hund $J$ values are derived [5]

$$\chi^{ij} = \left( \chi_{0}^{-1} - \chi^{-1} \right)_{ij}$$

Derivatives of the $\chi^{-1}$ matrix with respect to individual $\chi_{kl}$ can be obtained by the following relation:

$$\frac{\partial}{\partial \chi_{kl}} \left( \chi^{-1} \right) = -\chi^{-1} \left( \frac{\partial}{\partial \chi_{kl}} \chi \right) \chi^{-1}$$

where

$$\frac{\partial}{\partial \chi_{kl}} \{ \chi \}_{ij} = \begin{cases} 1 & \text{if } kl = ij \\ 0 & \text{otherwise} \end{cases}$$

(B1)

Using this fact, it is possible to obtain the full Jacobian of $f$ with respect to response $\chi$ matrices which can be used to obtain the covariance uncertainty matrix associated with the elements of $f_{ij}$, to a first-order expansion of $f_{ij}$ [74]

$$\Sigma_f = J_{\chi_{ij}} \Sigma_{\chi_{ij}} J_{\chi_{ij}}^T + J_{\chi} \Sigma_{\chi} J_{\chi}^T$$

(B2)
individual elements of χ and χ₀ are independent, we can assume that Σ covariance matrices are diagonal in order to make the following simplification:

\[
\sigma^2(f_{ij}) = \sum_{kl} \left( \frac{\partial}{\partial (\{\chi_0\})_{kl}} f_{ij} \right)^2 \sigma^2\left(\{\chi_0\}_{kl}\right) + \sum_{kl} \left( \frac{\partial}{\partial (\chi)_{kl}} f_{ij} \right)^2 \sigma^2(\chi_{kl}), \quad (B3)
\]

where \(\sigma^2(f_{ij}), \sigma^2(\{\chi_0\}_{ij}),\) and \(\sigma^2(\{\chi\}_{ij})\) correspond to the diagonal elements of \(\Sigma_f, \Sigma_{\chi_0},\) and \(\Sigma_{\chi},\) respectively.

With the established expression for the uncertainty values of \(f\) in Equation B3, we can express the squared uncertainty of \(U\), for an atomic site \(\gamma\), in the next level of uncertainty propagation,

\[
\sigma^2(U_{\gamma}) = \sum_{\sigma, \sigma'} \left( \frac{\partial}{\partial f_{\gamma\gamma}} G_U(f_{\gamma\gamma}) \right)^2 \sigma^2(f_{\sigma\sigma'}), \quad (B4)
\]

Equation B4 can be extended to an expression of the squared uncertainty of Hund \(J\), where \(G_U\) and \(G_J\) are functions of 2×2 sub-matrices along the diagonal of \(f\), as introduced in Equation 11, and depend on the different scaling schemes introduced in Ref. 5.

### Appendix C: Details of the data behind the periodic tables

#### TABLE V: The mean and standard deviation (σ) in the \(U\) and \(J\) parameters used in the periodic tables of Figure 1, alongside the number of samples \(N\).

| element | mean \(U\) | σ \(U\) | \(N\) | mean \(J\) | σ \(J\) | \(N\) |
|---------|------------|-------|-----|------------|-------|-----|
| Mn      | 4.710      | 0.707 | 94  | 0.575      | 0.157 | 97  |
| Fe      | 4.545      | 0.674 | 78  | 0.437      | 1.137 | 122 |
| V       | 3.909      | 0.404 | 68  | 0.849      | 0.538 | 108 |
| Cu      | 7.590      | 0.728 | 51  | 1.117      | 1.083 | 71  |
| Cr      | 2.982      | 0.464 | 47  | 0.557      | 0.089 | 61  |
| Nb      | 0.529      | 0.107 | 47  | 0.193      | 0.054 | 39  |
| Ti      | 4.737      | 0.428 | 45  | 0.705      | 0.861 | 62  |
| Ta      | 3.688      | 0.130 | 34  | 0.628      | 0.079 | 37  |
| W       | 1.846      | 0.213 | 33  | 0.385      | 0.045 | 33  |
| O       | 5.237      | 0.566 | 33  | 0.807      | 1.361 | 46  |
| Ag      | 2.830      | 0.606 | 26  | 0.703      | 0.131 | 24  |
| Re      | 0.598      | 0.172 | 26  | 0.255      | 0.089 | 27  |
| Ni      | 5.847      | 0.704 | 25  | 0.589      | 0.320 | 33  |
| Zr      | 4.382      | 0.269 | 24  | 0.740      | 0.069 | 23  |
| Mo      | 2.431      | 0.230 | 21  | 0.520      | 0.220 | 28  |
| Hg      | 0.520      | 0.220 | 21  | 0.288      | 0.271 | 22  |
| Cd      | 0.350      | 0.327 | 19  | 0.609      | 0.614 | 5   |
| Sc      | 2.506      | 0.210 | 16  | 0.543      | 0.070 | 16  |
| Y       | 4.704      | 0.393 | 15  | 0.825      | 0.179 | 5   |
| Pt      | 1.673      | 0.318 | 13  | 0.322      | 0.201 | 14  |
| Os      | 1.855      | 0.448 | 10  | 0.361      | 0.087 | 11  |
| Ru      | 2.972      | 0.548 | 10  | 0.504      | 0.292 | 24  |
| Lu      | 0.449      | 0.065 | 9   | 0.292      | 0.072 | 8   |
| Pd      | 3.608      | 0.407 | 8   | 0.620      | 0.106 | 10  |
| Hf      | 3.733      | 0.299 | 8   | 0.812      | 0.122 | 8   |
| Au      | 1.186      | 0.256 | 7   | 0.484      | 0.165 | 8   |
| Zn      | 0.530      | 0.795 | 5   | -0.105     | 0.433 | 17  |
| Rh      | 1.616      | 0.201 | 5   | 0.406      | 0.065 | 5   |
| Ir      | 1.868      | 0.288 | 5   | 0.352      | 0.283 | 14  |
| Tc      | 2.956      | 0.100 | 3   | 0.980      | 1.247 | 9   |
| Total   | 810        | 987   |     |            |       |     |

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