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Charge states of ions, and mechanisms of charge ordering transitions

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Abstract
To gain insight into the mechanism of charge ordering transitions, which conventionally are pictured as a disproportionation of an ion M as 2M⁺⁺⁺ → M⁺⁺⁺⁺ + M⁺⁺⁺⁻, we (1) review and reconsider the charge state (or oxidation number) picture itself, (2) introduce new results for the putative charge ordering compound AgNiO₂ and the dual charge state insulator AgO, and (3) analyze the cationic occupations of the actual (not formal) charge, and work to reconcile the conundrums that arise. We establish that several of the clearest cases of charge ordering transitions involve no disproportion (no charge transfer between the cations, and hence no charge ordering), and that the experimental data used to support charge ordering can be accounted for within density functional-based calculations that contain no charge transfer between cations. We propose that the charge state picture retains meaning and importance, at least in many cases, if one focuses on Wannier functions rather than atomic orbitals. The challenge of modeling charge ordering transitions with model Hamiltonians is discussed.

Keywords: charge state, oxidation state, charge order transition, correlation, transition metal oxide

(Some figures may appear in colour only in the online journal)

1. Introduction to the issues

As the momentous transition between conducting and insulating states by making a simple change (temperature, pressure, electron concentration, disorder), the metal–insulator transition (MIT) has assumed extraordinary importance in condensed matter physics research, and it plays a role in actual and anticipated applications. Established mechanisms include (1) interelectronic repulsion of electrons at low density (Wigner transition), (2) intratonic repulsion and resulting correlation (Mott transition), (3) disorder leading to incoherence and localization (Anderson transition), (4) coupling to the lattice, thereby opening new bandgaps (Peierls transition), and (5) interatomic repulsion leading to charge ordering (‘Verwey transition?’). The latter mechanism is distinguished by a breaking of symmetry on a sublattice of cations having partially filled correlated (namely 3d) electronic shells.

The mechanism for the Verwey MIT in magnetite Fe₃O₄ from the time of its discovery [1] was speculated to involve ordering to two charge states Fe³⁺ and Fe²⁺ on one Fe sublattice, making it the earliest example of a charge order transition, one that evidently would explain the MIT. A related charge order mechanism arises from disproportionation occurring at, and driving, the transition, which might also be envisioned as the mechanism in magnetite: 2Fe²⁺⁻⁻→Fe²⁺⁺ + Fe³⁺. The former is a disorder–order (charge disorder–charge order) transition in which the ‘symmetric’ metallic phase contains two space- and time-fluctuating charge states, and the second is a disproportionation-driven transition in which every site in the metallic phase is equivalent even on a short time scale ³ (rather than only in time-averaged diffraction).

³ The distinction may be one of the time scale of electron hopping between Fe sites, which will be reflected in the magnitude of the resistivity in the conducting phase.
In this paper, we extend our earlier work [2] on analyzing the connection between putative charge order driven MITs and the cation charge states that are involved. This study brings the concept, and the specification, of charge state or (formal) oxidation state or formal valence to the fore. We noted that the physical cation charge—the 3d occupation—can be identified from electronic structure calculations, by giving up the usual approach of integrating the density over some volume and instead simply looking at the cation charge density in the vicinity of the peak in the radial density $4\pi r^2 \rho(r)$ where only 3d density resides. Differences in 3d occupation, which are our current interest, are particularly easy to identify and quantify. Several examples of differing charge states were shown to contain equal 3d occupations; the actual charge is effectively divorced from the ‘charge state’ in some of those examples.

In section 2, the original chemist’s concept of ‘oxidation number’ (or (formal) oxidation state) is discussed briefly, noting that it metamorphosed into a more physical picture of ‘charge state’ for materials physicists. A brief poor man’s description of what is used to specify the charge state of an ion is provided in section 3. Some theoretical aspects, relating both to charge order transitions and to oxidation states, are discussed in section 4. In section 5, we begin by recounting one illustrative example of competing charge states in a compound, and proceed to review and extend slightly our previous examples of charge ordering materials where it was established that no 3d occupation difference between differing charge states exists. We proceed to an unconventional case—metallic with second-order transition—‘charge ordering’ compound AgNiO$_2$ in section 6, and in section 7 we look into the charge states of Ag in stable, non-transforming, insulating AgO. Our discussion in section 8 pulls together some of our thoughts on understanding and modeling structural and electronic transitions in the types of systems we have discussed.

2. Oxidation number

Chemists introduced the oxidation number (sometimes, oxidation state) of an atom in a molecule almost entirely on the basis of structure and electronegativity. With its high electronegativity, oxygen has oxidation number −2 except in very unusual geometries, and halides are −1, in both cases filling a shell. Electropositive ions (alkalis, alkaline earths, rare earths) in oxides and halides donate the electrons in their outer s and p shells, giving obvious positive oxidation numbers denoted early on and occasionally still by Roman numerals: namely Li$^I$, Ca$^{II}$, La$^{III}$. Most of the complications arise when, after nature has completed its bonding, an open d shell remains. For example, vanadium oxides may have V$^{IV}$, V$^{III}$, V$^{V}$, and V$^{IV}$, and combinations thereof.

The chemistry literature [3–6] contains numerous disclaimers that these oxidation states ‘have nothing to do with actual charge’, and some strive to maintain distinctions between the different terms in covalent molecules and molecular complexes. In the current materials physics literature there is no discernible distinction. Oxidation states are assigned from the structure (coordination and distance of neighbors of known valence), becoming formalized [7] in the valence bond sum that derives a formal valence (or ‘bond order’) directly and solely, from bond lengths. Sometimes the magnetic moment is involved, giving different ionic radii for high-spin and low-spin ions. Nevertheless, oxidation numbers (formal valences) have come to be connected with the number of actively occupied local orbitals; through size—higher oxidation state means fewer electrons and a smaller ion; through symmetry—a Jahn–Teller (JT) distorted coordination shell reflects specific orbital occupations; and through magnetic moment, which requires specific unpaired spin orbitals.

These oxidation numbers smelled a lot like charges to later scientists (especially material physicists), and they necessarily sum to zero like real charges. Already in 1939, 14 years after the naissance of quantum mechanics, Verwey was discussing the metal-to-insulator (MIT) transition in magnetite (Fe$_3$O$_4$)—the Verwey transition at 120 K—in terms of charge ordering of Fe$^{II}$ and Fe$^{III}$ ions on one of the iron sublattices. Note that the notation had already begun to shift to Fe(II) → Fe$^{II}+$, etc. The meaning was clearly physical charges as opposed to puerile oxidation states: to account for the transition from itinerant conductor to charge-localized insulator, actual charges were ordering, getting stuck to ions, becoming localized, and so forth—that seemed to be the problem to solve.

3. Assigning charge states of d ions

Ionic radii and high spin versus low spin designations differ with coordination [8], due in large part to the difference in crystal field. Unless noted, we use language applying to octahedral coordination for our comments.

Size. The ionic radius characterization of charge states has organized a great deal of structural information. The organization of all the structural data culminated in the table of Shannon ionic radii compiled by Shannon and Prewitt that has provided the standard for more than four decades [8]. For d ions in sixfold coordination, the average cation–oxygen distance together with the O$^{2−}$ radius of ∼1.40–1.42 Å specifies the ionic radius, the Shannon values being the best fit for a given presumed charge state to a large set of compounds. Note that the ionic radius contains no input from the actual size of the electronic cloud of the ion. Also, ionic radii differ for different coordinations, whereas the charge density is virtually invariant. The charge density is not used, as it is never measured with sufficient precision to be useful in this context (and chemists insist that it is irrelevant anyway).

Symmetry. Distortions from symmetric coordination—Jahn–Teller distortions—are assigned to the occupation of specific directional orbitals. Such assignments have been borne out by electronic structure calculations in a great number of cases. The La$_2$VCuO$_4$ example will be discussed in section 5, where it will be seen that spin-orbital occupations can be confusing.

Magnetic moment. A Curie–Weiss moment, or ordered moment, reflects ‘unpaired’ occupation, i.e. distinct orbitals are occupied by one spin only. Cations of a given charge state can sometimes occur in both high-spin and low-spin
(often zero) configurations. Interestingly, the two spin states that arise in several ions have been assigned to different ionic radii, with the high-spin configuration typically 0.04 Å larger than the low-spin configuration of a given charge state [8]. Spin polarization actually changes the 3d occupation and the radial density (real ionic size) by a negligible amount. Spin polarization does however create a spin splitting so that up and down spin electrons hybridize differently with the O ions, and there is an interorbital Hund’s rule energy.

4. Theoretical notions

4.1. Modeling of charge order transitions

Tight binding modeling of charge ordering transitions typically invokes, beyond the hopping term $H_K$, direct intersite repulsion between charges on neighboring cations in addition to intrasite repulsion:

$$H_U + H_V = U \sum_j n_{j,s} n_{j,-s} + \sum_{(ij),ss'} V_{ij} n_{i,s} n_{j,s'},$$  \(1\)

where $U$ is the Hubbard on-site repulsion and $V_{ij}$ is an intersite repulsive energy, where $(ij)$ indicates neighboring sites and $s, s'$ are spin indices. The second term alone, which is pictured as the driving force, would minimize the energy by forcing as much charge as possible onto as few distant sites as possible. The Hubbard term provides the necessary, and physical, energy that opposes excessive inhomogeneity. Pietig et al [9] and Hellberg [10] have shown, for example, that this Hamiltonian alone, on a two-dimensional cluster with periodic boundary conditions, shows a range of re-entrant (with temperature) charge order transitions in a certain range of $V/U$ (with $U$ set equal to the bandwidth), similar to the popular interpretation of behavior observed in some manganites.

Other Hamiltonian terms are included in more general models than the single-band case just mentioned: very often Jahn–Teller distortion energies $H_{JT}$ for multiorbital systems, and sometimes magnetic Hund’s rule energy ($on-site$ spin–spin) $H_{HR}$ as well as distance-dependent kinetic hopping $H_K$. This type of extension (without Hund’s rule) has been applied by Mishra et al [11] to model behavior observed in manganites. In a two-band model for $e_g$ electrons in perovskite nickelates (which we discuss below), Lee and coworkers [12] applied a Hamiltonian $H_K + H_U + H_{HR}$ (notably without $H_V$) and obtained charge order-like phases driven by Fermi surface nesting. These nickelates do not show JT distortions as anticipated of an $e_g$ system, prompting Mazin et al [13] to observe that avoidance of JT distortion can be accomplished in certain regimes by charge ordering instead. They suggested that the form of Hamiltonian mentioned above, minus the $H_V$ term, should provide the relevant energies. Interestingly also, when performing a thorough mean-field study of ordered states in ANiO$_2$ ($A =$ Na, Li, Ag, the latter of which we address in section 6), Uchigaito et al also chose not to use the $H_V$ intersite repulsion term, although several other interactions were included. Amongst the many possible phases uncovered for a selection of independent interaction strengths in the Hamiltonian were phases of charge-ordered type [14].

4.2. Identification of charge states/oxidation states

It is commonly stated and generally accepted that dividing the crystalline density into atomic contributions is so subjective as to be useless for specifying a charge state. This statement contains a lot of truth but is not entirely correct. Creating disjoint volumes associated with ions is indeed subjective and ambiguous. However, the general question is more subtle than dividing up space, so it is worthwhile to digress briefly.

4.2.1. From pseudoatoms to the enatom. Almost four decades ago Ball demonstrated [15], using small displacements of nuclei, how the charge density of a collection of atoms can be decomposed uniquely into contributions from each atom (‘pseudoatoms’). In the process he also identified a charge backflow field associated with each atom when it is displaced slightly. Subsequently he qualified [16] this result, which does indeed apply to any finite system and to non-polar solids where the pseudoatoms are necessarily neutral. Polar solids, however, provide additional challenges closely related to the difficulties of treating polarization in such solids, where much progress has been made in the past two decades [17, 18]. The fact that the Born effective charge for a charged ion is a tensor provides insight: the (Born) effective charge is not a simple scalar, but depends on the direction of displacement. Ball’s prescription has so far been applied only to simple metals [19], where it may be useful for understanding lattice dynamics and, applying the same prescription to the potential, electron–phonon coupling [20]. Because the name pseudoatom has been used in so many different contexts, the designation enatom has been suggested for Ball’s pseudoatom [19].

It remains the case that trying to divide space into regions in which the charge density is assigned to a particular atom is a subjective enterprise. Ball’s approach differs, by deriving overlapping atomic densities extending perhaps to a few shells of neighbors. This observation conforms well to our finding [2], discussed at more length below, that the d occupation often does not vary at all between different charge states. If a rather small sphere were chosen to specify the charge, different charge states would have identical charges (in those regions). Where the charge state picture clearly works—which is very often—the electronic spectrum is a central property: occupied bands corresponding to occupied local spin orbitals can be identified, and the sum of the orbital occupations determines the charge state. These have conventionally been pictured as atomic orbitals, but the development and use of Wannier functions (WFs) over the past decade provides an alternative viewpoint that we return to below.

A well-specified division of space into regions associated with atoms is provided by the Bader prescription [21]. The ‘zero flux’ (of density) surface is defined as that surface for which the density is a minimum over (small) distances perpendicular to the surface; thus it depends solely on the density $\rho(\vec{r})$. While probably the most reasonable ‘division into atom-based regions’ of space, it has evident peculiarities, at least in details. Consider a density comprised simply of overlapping neutral-atom densities, so each atom is neutral by supposition. The resulting Bader charge will not be zero...
and will depend on the distance between atoms (i.e. on the environment). Values of Bader charge have not been reported extensively in the solid state literature (more so for molecules); however the capability has been implemented in some electronic structure codes, and we provide Bader charge values in section 8 for some of the charge ordering systems we discuss in this paper.

4.2.2. Calculation of the oxidation state. Two proposals for the definition and calculation of ‘oxidation states’ (both used this terminology rather than charge states) have appeared recently. With molecular complexes in mind, Sit et al suggested [22] projecting the occupied states onto atomic (3d, say) orbitals to obtain the spin-orbital occupation matrix as is done in the LDA+U method, to diagonalize this matrix. The number of eigenvalues representing full occupancy then provides the number of occupied physical orbitals and hence the oxidation state. Smaller occupations are ignored. For the transition metal complexes they provided as examples, the procedure was relatively clear. However, possible ambiguities could be imagined: in one of the examples 0.93 needed to be interpreted as occupied, 0.63 as not occupied. More testing of these ideas should prove instructive.

Another proposal, by Jiang, Levchenko, and Rappe [23], applying specifically to periodic systems, employed developments in the theory of polarization in crystals to formulate their definition of oxidation number as the integral (of namely the average of) the Born effective charge as the ion’s sublattice is displaced adiabatically by one direct lattice vector. The material must remain insulating, but otherwise the result is independent of the path in configuration space. This procedure does not identify an ion’s charge density or necessarily even entail a true charge, but it does provide the interpretation of the change in polarization as being due to the transport of a number of unit charges equal to the oxidation state through a displacement corresponding to the lattice vector.

As in the case of the enatom mentioned earlier, this definition also does not make the oxidation state a property of the static reference state—displacement is required. Because the path is periodic (it can be repeated), any point can be considered as the starting (reference) point, with the same oxidation state (same integral). Thus the ion retains one fixed oxidation state along the path in configuration space. Considering that the Born effective charge is anisotropic, it seems possible that the oxidation state by this definition in an anisotropic insulator also depends—as does the Born effective charge—on the direction of the direct lattice vector that connects the end points [24]. Further applications of this proposal will be enlightening, as will the fact that displacement is required to specify what is normally envisioned as a property of the static system.

4.2.3. Oxidation states from Wannier functions. In the survey in this paper of charge ordering transition involving different charge states, and hence of the specification of charge states more generally, we will begin to explore the viability of abandoning atomic orbitals, whose occupation often does not change at the ‘charge ordering’ transition, in favor of Wannier orbitals. The basic idea is as follows: calculate atomic-like Wannier functions for the relevant states and explore whether these orbitals support the charge state viewpoint. Ideas along these lines have been implicit in discussing specific materials, and for some symmetric and therefore more straightforward cases this picture has been stated explicitly, most recently by Haverkort et al [25]. Wannier functions have been integrated into the electronic structure calculations in various ways, so this change from atomic to Wannier orbitals becomes transformational when applied to ‘charge ordering’ transitions. Without any change in actual (3d) charge on the open shell cation, what are the relevant degrees of freedom to consider? We return to some of these general issues in section 8, but we begin by considering a simple and nearly transparent case illustrating the idea.

In the double perovskite compound Ba$_2$NaOsO$_6$, osmium is formally a heptavalent Os$^{7+}$ 5d$^1$ ion—a very high oxidation state. Calculating the 5d charge in the ion, though one cannot arrive at a precise value as noted above, corresponds to occupation by around 4.5 (not 1.0) 5d electrons [26]. Cases of large differences between the actual charge and the oxidation state are common knowledge in the electronic structure community. Does this signal the breakdown of the charge state picture for high valence state ions? It does not: calculations [27, 28] indicate one electron in the otherwise unoccupied ‘5d bands’ spins half of the course, and if the 5d occupation is not tied to the charge state (which it is not) then the formal valence picture survives in robust fashion.

Analysis shows that the occupied 5d orbital (i.e. the Wannier function) is a strongly hybridized mixture of 50% Os 5d state and 50% O 2p contributions distributed around the neighboring oxygen ions. The other four electrons making up the total 5d occupation of 4.5 are hybridized into (and thereby vanish into) the 6 atoms × 3 orbitals = 18 occupied O ‘2p bands’. Transformation from atomic orbitals to Wannier orbitals would presumably lead to 18 occupied O-centered orbitals of p symmetry with tails of Os 5d states, and one occupied Os 5d-derived orbital with tails composed of O 2p orbitals. So although several 5d electrons vanish into the O 2p bands, all five ‘5d orbitals’ survive by recovering charge from the 2p orbitals. Understanding that strong metal–oxygen hybridization implies the efficacy of generalized (Wannier) orbitals has been around for a while; for example, it provided a simple picture explaining why second Cu–Cu neighbor coupling along a chain dominated over near-neighbor coupling in the Cu–O magnetic chain compound Li$_2$CuO [29].

Another example illustrates an apparent connection between WFs and formal valence. Volja et al [30] have calculated $e_g$ WFs for the two charge states in a charge-ordered manganite La$_{1/2}$Ca$_{1/2}$MnO$_3$, finding a substantial difference in the extent of the tails, the ‘Mn$^{4+}$’ WF being more localized. Concerns about specifying charge states with WFs include (1) a computational issue: the gauge freedom means that WFs are far from unique, so subjective choices may lead to different conclusions, and (2) a conceptual issue: the choice of projections—the orientations, for example—they themselves are not objective. At one extreme are maximally localized WFs [31, 32] that may not be centered on an ion and may
be unsymmetrical, therefore complicating if not destroying any utility in representing physical orbitals. Atomic-orbital and symmetry projected WFs [33] (‘maximally projected’) provide an alternative approach, one that should provide a more physical representation, and the projection issue may present less of a non-uniqueness ambiguity. This viewpoint remains to be tested seriously.

5. 3d transition metal charge states: illustrative examples

In oxides containing two transition metals, one of the first tasks in understanding them is to categorize the charge states of the ions. The sum is known from oxygen and the simpler ions, but a number of competing factors (which are not well understood and are likely to be material specific) determine the balance. The double perovskite structure is simple, and it provides our first example.

5.1. Competing charge states of La₆VCuO₆

The computational methods we have used have been described previously [2, 34]. In our earlier work [35] on La₂VCuO₆, using the LDA+U method [36, 37] as is necessary for such oxides, it was found that the two configurations V⁺⁺Cu¹⁺, a standard closed-shell band insulator, and V⁺⁺Cu²⁺, a spin-compensated ferrimagnetic correlated insulator with a very small gap, are practically degenerate in energy after full structural relaxation. These charge state assignments fit in all respects. Structurally, the mean V–O distances are 1.84 Å and 1.88 Å, respectively, for the two configurations, differing by the difference in the corresponding V ionic radii (0.04 Å). The mean Cu–O distances are 2.02 Å and 1.98 Å, respectively, again differing by their ionic radii (0.04 Å also). The Jahn–Teller distortion magnitude for the magnetic configuration (long bond minus short bond) is 0.11 Å for both ions, an unambiguous JT distortion. Magnetically, V and Cu have one electron and one hole, respectively, as judged by the moment (reduced somewhat by hybridization) and by their respective filled and empty bands. These two configurations are ‘poster kids’ for the charge state picture—nothing could be clearer. The spin density is pictured in figure 1 and supports the charge state description of the spin-compensated Mott insulating phase.

What our earlier work [2] also established is that the total 3d occupation in both charge states of V, and also of Cu, are identical (by this we mean 0.5% difference or less—no physically meaningful difference). It has occasionally been noted in the literature of electronic structure calculations (including our own) that atomic charges in different charge states ‘differ rather little’ in charge. In this case, and in the few others that we have checked so far (see below), the difference is negligible.

4 The density functional-based linearized augmented plane wave code wien2k [34] has been used in our calculations, where we have applied correlation corrections (LDA+U) as is usually necessary to describe 3d oxides. Computational methods are as described in [35].
are identical for the two charge states. Values for the active Jahn–Teller orbitals are emphasized in boldface.

The difference in the total (last column) for the two charge states of both V and Cu arise from 'charge state' obscures the issue rather than clarifying it. It is this conundrum that we will expand on in the following sections. Note at this point that we have tried not to say ‘what local orbitals are occupied’ but instead what local spin environments. Using the term ‘oxidation state’ instead of ‘charge state’ is an essential concept in conveying the character of the ion and its environs. Yet charge state very often is an oxymoron, speciﬁc a specific charge seems to classify it as an oxymoron, yet charge state very often is an essential concept in conveying the character of the ion and its environs (‘which local spin orbitals are occupied’). It is this conundrum that we will expand on in the following sections. Note at this point that we have tried not to say ‘what atomic orbitals are occupied’ but instead what local orbitals are occupied. We provide most of our discussion that follows, with one exception, in the context of materials with ‘charge ordering transitions’.

5.2. Rare earth nickelates

The structural and metal–insulator transition [38, 39] in the rare earth nickelates RNiO	extsubscript{3} has been discussed for two decades as a charge order transition, with experimental data being analyzed in considerable detail for YNiO	extsubscript{3} and several others in terms of a fractional charge transfer 2Ni	extsuperscript{3+} → Ni	extsuperscript{3+} + Ni	extsuperscript{3–}, with the latest analysis [40] pointing to δ ≈ 0.3 for YNiO	extsubscript{3}. The high-temperature diffraction-determined structure has a single Ni site in the distorted GdFeO	extsubscript{3}-type cell containing four formula units with a single Ni site, while two separate sites Ni	extsubscript{1} and Ni	extsubscript{2} appear in the low-temperature symmetry-broken (charge-ordered) phase. The Ni–O octahedra remain nearly regular (i.e. without JT distortion), with the Ni–O and Ni2–O mean distances being 2.02 Å and 1.92 Å, respectively. A simple breathing mode mediates the transition. The large Ni–O separation differences are consistent with the Ni	extsuperscript{2+} and Ni	extsuperscript{4+} charge state picture, except for δ = 1 rather than δ = 0.3. There are, however, several questions about the current interpretation that we now discuss.

Our investigation into mechanisms of ‘charge ordering’ transitions was spurred by noticing [2] that the Ni	extsubscript{1} and Ni	extsubscript{2} sites have identical 3d charge, being also the same as in the high-symmetry phase with a single Ni site. In figure 2, a different type of presentation is provided. A single ‘Ni	extsuperscript{2+}’ radial density is plotted, that of one of the sites in AgNiO	extsubscript{2} (see below); the other Ni ions in this figure, even those with different charge states, are indistinguishable on this scale in the region of the 3d peak. With this ‘Ni	extsuperscript{2+}’ site as the reference, the relative differences (expressed in percentages) for ‘Ni	extsuperscript{(3–δ)+}, (Ni	extsubscript{1} in YNiO	extsubscript{2}), ‘Ni	extsuperscript{(3–δ)+}, (Ni	extsubscript{2} in YNiO	extsubscript{3}), and ‘Ni	extsuperscript{3+}, (Ni	extsubscript{1} in AgNiO	extsubscript{2}) are plotted. In the region of the peak in density r = 0.6a	extsubscript{o}, the differences are of the order of 0.5%. The shapes of the density (namely 3d orbital) become slightly different around 1a	extsubscript{o}, and the effects of oxygen tails at different distances become evident beyond 1.4a	extsubscript{o}. Note that the ‘Ni	extsuperscript{3+}’ density is the nearer to the ‘Ni	extsuperscript{2+}’ density than are the other two.

Since the NiO	extsubscript{6} octahedra are somewhat distorted in this system, it is the eigenvalues of the occupation matrices for the two spins for (ferromagnetic) YNiO	extsubscript{3} that we present in table 2. These values can be used to evaluate the oxidation state prescription of Sit et al. [22]. Their prescription gives Ni	extsuperscript{2+} and Ni	extsuperscript{3+} oxidation states as long as one identifies 0.83 and 0.85 as ‘fully occupied’ (the other fully occupied states have eigenvalue 0.91–0.92), and 0.58 and 0.64 as not fully occupied and thus unoccupied. To make the issue more apparent, it is

<table>
<thead>
<tr>
<th>Atom</th>
<th>xy</th>
<th>xz</th>
<th>yz</th>
<th>x² − y²</th>
<th>3z² − r²</th>
<th>Total</th>
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<td>V⁴⁺:d⁴ up</td>
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<td>0.11</td>
<td>0.26</td>
<td>0.18</td>
<td>1.40</td>
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<tr>
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<td>0.26</td>
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<tr>
<td>V⁴⁺:d⁴ diff</td>
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<td>0.01</td>
<td>0.00</td>
<td>0.00</td>
<td>0.72</td>
</tr>
<tr>
<td>V⁴⁺:d⁴ sum</td>
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<td>0.20</td>
<td>0.20</td>
<td>0.51</td>
<td>0.36</td>
<td>2.11</td>
</tr>
<tr>
<td>V⁵⁺:d⁴</td>
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<td>0.30</td>
<td>0.30</td>
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<td>0.57</td>
<td>2.04</td>
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<tr>
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<td>0.95</td>
<td>0.29</td>
<td>4.03</td>
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<tr>
<td>Cu²⁺:d⁹ dn</td>
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<tr>
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<td>1.75</td>
<td>1.86</td>
<td>1.86</td>
<td>8.97</td>
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</tbody>
</table>

Figure 2. Plot (dashed line) of the radial density of the so-called ‘Ni	extsuperscript{2+}’ ion (Ni1) in AgNiO	extsubscript{2}, together with the percentage differences (with this ion as the reference) of the other Ni sites noted in the legend. In the region of the peak the differences are 0.5%, even between different compounds. The differences that become large in the tail region illustrate the sensitivity in this region to the environment of O 2p orbital tails.
Table 2. Eigenvalues of the spin-orbital occupation matrices for the Ni1 (1.35 µB) and Ni2 (0.55 µB) sites in ‘charge-ordered’ YNiO3, from a ferromagnetic LDA+U calculation. Values for ‘non-filled orbitals’ are emphasized in boldface. Notice the absence of appreciable Jahn–Teller splitting (difference between occupations of \(x^2−y^2\) and \(3z^2−r^2\)).

<table>
<thead>
<tr>
<th>Eigenvalue</th>
<th>(xy)</th>
<th>(zx)</th>
<th>(yz)</th>
<th>(x^2−y^2)</th>
<th>(3z^2−r^2)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni1 up</td>
<td>0.91</td>
<td>0.92</td>
<td>0.92</td>
<td>0.85</td>
<td>0.83</td>
<td>4.43</td>
</tr>
<tr>
<td>Ni1 dn</td>
<td>0.91</td>
<td>0.91</td>
<td>0.91</td>
<td>0.19</td>
<td>0.17</td>
<td>3.08</td>
</tr>
<tr>
<td>Ni2 up</td>
<td>0.92</td>
<td>0.92</td>
<td>0.92</td>
<td>0.64</td>
<td>0.58</td>
<td>3.98</td>
</tr>
<tr>
<td>Ni2 dn</td>
<td>0.92</td>
<td>0.92</td>
<td>0.92</td>
<td>0.36</td>
<td>0.32</td>
<td>3.44</td>
</tr>
</tbody>
</table>

useful to think of scaling the maximum occupation for this sphere size, 0.92, up to 1.00. Then 0.84 → 0.91 must be ‘fully occupied’ and 0.64 → 0.70 must be treated as ‘not fully occupied’ to give the mentioned oxidation states. The atomic moments are calculated as 1.35 µB and 0.54µB versus the idealized values of 2 and 0 µB, respectively.

The atomic-orbital projected density of states (PDOS), presented earlier [2, 13], is reproduced in figure 3. The larger moment on Ni1 arises almost equally from more majority states centered at −0.5 eV (an expected difference, near \(E_F\)) and fewer minority states near −5 eV (an unexpected change in the strongly bound part of the spectrum). The amount of unoccupied 3d spectral density does not differ much between Ni1 and Ni2, unlike the formal picture would suggest (a factor of 2) but it is consistent with our finding of no difference of 3d occupation. The difference lies in the spin and energetic distribution. The ‘charge state’ picture thus is murky; the ionic radii are consistent with full disproportionation \(\delta = 1\), whereas other data (see below) are used to justify \(\delta = 0.3\) as the best value. The spin-orbital occupation eigenvalues and the PDOS are hardly definitive.

Very often the contrasting core-level energies of different sites are used to identify and specify charge states, and this has been done for a few nickelates. In our earlier paper [2], we pointed out that these on-site energies are given well by density functional theory (DFT) calculations, and thus are due to the differing environments rather than to any difference in local charge.

5.2.1. Charge order versus Jahn–Teller distortion. In charge ordering transition phenomenology, or more broadly for the specification of the oxidation state, the real cation charge—the 3d occupancy—is a peripheral (many say irrelevant, as mentioned above) issue, and our specification of 3d occupation reinforces that picture. However, the charge state language invites—in essence, requires—an interpretation of selected spin orbitals to account for differences in the different sites. The ordered state of RNiO3, which contains roughly symmetric NiO6 octahedra of substantially different size, cannot be understood in terms of an \(e^1_g\) ion which should have strong JT tendencies; hence the conventional ‘charge order’ designation. This picture would have the driving mechanism, the entropy frozen out at the transition, due to charges hopping amongst disordered Ni2+ and Ni4+ sites.

Mazin et al [13] observed that a fully disproportionated ‘high-spin \(e^2_g\), no-spin \(e^0_g\) ’ picture accounts for (1) the lack of JT distortion and (2) the large and small Ni moments.

They confined their attention to the ordered phase (ground state). The large and small octahedra, expected on the basis of differences in ionic radii for distinct charge states, does not have any quantitative explanation—in any picture—for ions of identical charge and radial extent. They further suggested that the Hund’s rule magnetic energy is important in producing these configurations. Our earlier work established that there is negligible actual 3d occupation difference between Ni1 and Ni2, which seeks to leave the search for the true microscopic driving force between the charge transfer energy, Hund’s rule energy, and electron–lattice coupling, and the change in kinetic energy that results from the latter two and modulated by the former.

5.2.2. Disorder–order scenario. X-ray absorption spectra for nickelates suggest a very different view of the transition. Spectral signatures of the two distinct Ni sites are found in the high-temperature symmetric phase [41, 42] as well as beyond the phase transition under pressure [43]. The interpretation must be that the Ni1 and Ni2 sites are already different above the transition, but are disordered and fluctuating in both space and time. The ionic radii are not a good fit for the perovskite structure, and the structure adjusts by distorting to large and small octahedra. Without structural coherence, carriers can hop—the material is conducting although only as a very bad
metal. The entropy is due to fluctuating local breathing modes that freeze in at the transition.

In the ordered phase, the coherent alternation of distortions—a non-zero amplitude of the zone boundary breathing mode, to use common language—opens a gap in the band structure, and the material becomes a (correlated) insulator. The transition can be approached from the opposite direction: the ordered phase is a small-gap Mott insulator as long as the structural distortion is coherent, but loss of the coherence smears the gap and leads to a very bad metal but conducting phase. From either viewpoint, the transition is at the most basic level an order–disorder transition, rendered more complex than most by the electronic correlation effects.

The breathing distortion modulates the Ni 3d on-site energy, and hence modulates the charge transfer energy between the O subsystem and each Ni site. JT distortions, on the other hand, retain the mean Ni–O distance and do not modulate the 3d on-site energy significantly.

5.3. Results for other charge order systems

In our previous report [2], we noted two additional charge order systems for which there is no difference in 3d occupation of the ‘charge-ordered’ cations.

CaFeO$_2$. This ferrate is isostructural with the RNiO$_3$ class and displays a similar MIT and structural change. The proposed disproportionation [44] invokes the unusually high oxidation state Fe$^{5+}$ in addition to the charge-balancing Fe$^{3+}$ state. The 3d occupations, determined from the radial charge densities, in the charge-ordered states for the two Fe sites are identical. It seems plausible that the disorder–order scenario proposed for RNiO$_3$ applies to CaFeO$_2$ as well, though there is much less data to validate it. Yang and collaborators [45] noted from their DFT calculations that no meaningful (i.e. discernible) difference in real Fe 3d charge (which they identified as around 5.1 electrons) could be obtained, which led them to categorize the ordered state in terms of ligand holes, $d^5 + d^5L^2$, rather than as Fe$^{3+} + Fe^{5+}$. The difference in Fe–O distance changes the hybridization considerably, and the resulting difference in hyperfine field that they calculated agreed well with the experimental data.

V$_4$O$_7$. Vanadates of many stoichiometries display MITs accompanied by structural transitions, and the intricacies of the structures compound identification of the mechanism underlying the transitions. The low-temperature ordered phase of V$_4$O$_7$ has been characterized [46, 47] in terms of two distinct V$^{3+}$ sites and two V$^{4+}$ sites. Comparing the radial densities from DFT-based calculations [47], it became clear that all four of the V sites have the same 3d occupation, as for the various other systems we have surveyed and discussed in this paper. The difference in deep core levels for the two Fe sites is 0.9–1.2 eV, similar to the calculated and measured values for the nickelates and CaFeO$_3$, and used to substantiate ‘charge order’.

6. Charge order transition in AgNiO$_2$

The structural transition at 365 K in this triangular lattice compound is not first order but instead continuous, and moreover it is not an MIT but rather a (semi)metal to (semi)metal transition, so AgNiO$_2$ belongs to a separate class from the systems discussed above. It has nevertheless been characterized in some detail as a charge order system possessing other distinctions. It has been studied also for its unusual low-temperature magnetic behavior and ordering, which are not our primary concern. The high-temperature phase, while conducting, has a resistivity of 2–3 $\text{m} \Omega \text{cm}$, thus classifying it as a very bad metal or, more realistically, as a semimetal. Decreasing the temperature from the symmetric phase, the resistivity increases slightly but sharply below the transition before leveling off around $40^\circ$ below the transition and again resuming a positive (metallic) temperature derivative. This transport behavior is at a second-order structural transition, where the freezing out of lattice fluctuations upon ordering normally leads to a decrease in resistivity.

The charge state in the symmetric high-temperature phase, to the extent that one applies in a conductor, must be Ni$^{3+}$, so the triangular lattice of Ni sites provides a platform for frustration of orbital order as well as for potential magnetic order. The transition is to a $\sqrt{3} \times \sqrt{3}$ increase of the cell caused by oxygen displacement radially outward (within its basal plane) from one Ni, denoted Ni1, creating a large NiIO$_6$ octahedron and two smaller octahedra Ni2O$_6$ and Ni3O$_6$. Ni2 and Ni3 sites differ only due to the stacking sequence of NiO$_2$ slabs, and while this distinction is identifiable in some quantities, the difference is too small to be of interest here. At low temperature, Ni1 has a moment of $\sim 1.5 \mu_B$, while the moments on Ni2 and Ni3 are difficult to quantify, but suggested to be perhaps 0.1$\mu_B$. The Curie–Weiss moment, both above and for a wide temperature range below $T_c$, is roughly consistent with either a moment of 0.7–0.8$\mu_B$ on each Ni (the high-$T$ phase value), or 1.5$\mu_B$ on 1/3 of the Ni ions (the low-$T$ phase values). The data and the analysis are not precise enough to distinguish between these two possibilities, or something intermediate.

Wawrzynska et al pointed out [48, 49] that DFT calculations indicate that the Ni1 $e_g$ bands are split $\sim 1$ eV above and below $E_F$ (figure 4), giving it an $S = 1$ (Ni$^{2+}$) ‘Mott insulator’ configuration (even though no strong correlation corrections need to be used in the DFT calculations), with moment reduced by hybridization to be consistent with the observed value. Ni2 and Ni3 are only weakly magnetic, with their $e_g$ bands crossing $E_F$ (figure 4) consistent with an unpolarized ‘Ni$^{3.5+}$’ designation for both ions, necessitated by charge neutrality. The O displacement amplitude at low temperature is 0.06 Å, giving a change in Ni–O distances due to the distortion of $\sim 0.04$ Å. This amount is perhaps reasonable for oxidation states differing by 1/2, but Shannon’s Ni$^{2+} + Ni^{3+}$ difference is 0.11 Å, suggesting that a Ni$^{2+} + Ni^{3.5+}$ difference should be around 0.15 Å rather than 0.04 Å.

As we found for the other compounds which are said to display different charge states, we find that the 3d occupations in ‘charge-ordered’ AgNiO$_2$, from the radial charge densities
Figure 4. Projected DOS for $e_g$ and $t_{2g}$ orbitals for Ni1, Ni2, and Ni3 sites in ordered AgNiO$_2$. Majority is plotted upward, and minority downward. The spectral distributions of Ni2 and Ni3, with calculated moments of 0.28 and 0.27 $\mu_B$, respectively, show no discernible differences. The Ni1 moment of 1.27 $\mu_B$ is accompanied by the 2 eV $e_g$ spin splitting, shown at the top.

of the three Ni sites near their peaks, are indistinguishable—the 3d charge on each is the same. More detailed results on AgNiO$_2$, including energetics and magnetic moments during oxygen displacement, the Fermi surfaces in the paramagnetic state, and comparison with available experimental data, will be published elsewhere.

7. Charge states in AgO

Copper with its various charge states has been well studied because of its central importance in high-temperature superconductors. The isovalent 4d sister Ag rarely displays correlated behavior in ionic compounds or departs from its monovalent closed-shell configuration. The behavior observed in AgO which we now review puts it in an unusual borderline class of materials containing group IB cations.

AgO does not display any charge order or other transition, being insulating and diamagnetic at all temperatures. It has however been characterized as displaying both Ag$^+$ and Ag$^{3+}$ charge states. The crystal structure, pictured in figure 5, is monoclinic $P2_1/c$ (space group no. 14) [50, 51], $a = 5.86$ Å, $b = 3.48$ Å, $c = 5.50$ Å, $\beta = 107.5$, with two inequivalent Ag sites, as pictured in figure 5. The Ag1 (Wyckoff site 2d) sits at the center of a slightly distorted square, Ag2 (site 2a) forms a linear O–Ag2–O trimer of length 4.34 Å, and the O ion sits at the Wyckoff 4e site. The average Ag1–O bond length is 2.03 Å, while the Ag2–O separation is 2.17 Å, consistent with a lower charge state and its lower coordination. Each O (only one crystallographic site) connects two AgO$_4$ squares which are almost perpendicular and one O–Ag–O trimer. GGA and hybrid functional calculations by Allen et al were interpreted to support the two-charge-state picture: an unoccupied band is Ag1 derived (one band per Ag1 ion), while all of the Ag2 4d bands are occupied [52, 53].

To check the 4d occupation and do some further analysis, we carried out LDA+$U$ ($U = 5$ eV, $J = 0.68$ eV) calculations using the experimental structure. The atom-projected DOS (PDOS) is plotted in figure 6. The Ag1 PDOS dominates in the strongly bound region, from $-7.5$ to $-3.5$ eV, is smaller than that of Ag2 in the $-3.5$ to $-1.5$ eV, and then is nearly negligible (‘gapped’) from $-1.5$ eV through the bandgap. The unoccupied Ag1-derived band (one band per Ag1 ion) is, however, roughly half O 2p character, an occurrence that is sometimes characterized as ‘O 2p holes’. This latter picture would then assign the charge configuration Ag$^2+$L, where L is a ligand (oxygen) hole, instead of Ag$^{3+}$. Such a viewpoint becomes difficult to sustain, however, because the seemingly apparent Ag$^+$ charge state requires it to be surrounded by O$^{2-}$ ions, and there is only the single O ion coordinating with both Ag1 and Ag2.

The Ag2 PDOS is contained primarily in a 4 eV wide peak centered at $-2.5$ eV, plus the band just below the gap that
is half O 2p in character. The O 2p PDOS is spread relatively evenly through the range of the 4d bands, mixing strongly with both Ag1 and Ag2 except for some Ag2 bands in the −2 to −3.5 eV range.

As in the several cases mentioned above, we find negligible difference in the 4d occupation of actual charge (the radial charge densities are nearly identical near the peak). This negligible difference in 4d charge, as well as the extreme similarity to the Ag$^+$ ion density in AgNiO$_2$, is illustrated in figure 8.

The oxidation state specification of Sit et al can be tested here, using the natural local coordinate system where the orbital occupation matrix is nearly diagonal. For Ag2, four orbitals have occupation very near 0.80, i.e. nominal full occupation, since $\sim 20\%$ of the charge extends out of the LAPW sphere. The other, $d_z$, has occupation 0.75. Supposing that 94\% also corresponds to a filled orbital, Ag2 is indeed $d^{10}$, Ag$^2+$. For Ag1 in the square, four orbitals have $\sim 0.82$ occupation, while the $d_z$ orbital has occupation 0.54, about 68\% full. Sit et al’s prescription would be to ignore this, obtaining $d^8$Ag$^{3+}$.

A more illustrative picture is provided by the Ag1- and Ag2-projected fatbands plotted in figure 7, which illustrate several characteristics. Much of the weight of the Ag1 atom lies in the lower regions of the 4d bands, which are mixed strongly with O 2p states. A strong ligand field splitting separates one orbital—the $d_{z^2}$ member that antibonds strongly with the neighboring O 2p orbitals. The Ag2 $1^+$ ion in the trimer has most of its weight from $-4$ eV to $-1$ eV, with a split off antibonding $d_{z^2}-p_z$ combination forming a 1 eV narrow band below the gap.

8. Discussion

Briefly stated, we have observed that for several cases of ‘charge ordering’ in solids, usually involving a first-order metal–insulator transition, there is no change in cation 3d occupation across the transition, and hence no real charge ordering. This lack of any charge transfer between cations during a ‘charge ordering’ transition raises fundamental questions about the mechanism(s) behind these transitions and the best way to understand the underlying physics. There has been some discussion of a similar conundrum in magnetite by Garcia and Subias [54]. At a more basic level, this observation raises perhaps more basic questions about the meaning and specification of ‘charge states’ in solids in addition to the underlying mechanism—just what it is that orders at the transition; where is the entropy gained or lost?

Before continuing this discussion of charge, we present here the Bader charges $Q_B$ promised in section 1 for some of the systems discussed here. These values were obtained from the Wien2k code (based on the deviation from charge neutrality from the sum, the Bader charge is accurate to about 0.02/atom in magnitude).

- $YNiO_3$. Distorted phase: $Q_B$(Ni1) = +1.33, $Q_B$(Ni2) = +1.49, $Q_B$(Y) = +2.15. $Q_B$(O) = $-1.19 \pm 0.01$ for the three O sites. In the higher-symmetry phase, $Q_B$(Ni) = +1.45, with little change for Y and O. These values are from LDA+$U$ calculations with $U \approx 5.5$ eV. The Ni charges decrease somewhat with decrease of $U$, retaining about the same difference in the distorted phase. Compare with the formal values $+2$, +4, +3, and $-2$, respectively.

- $AgO$. Distorted phase: $Q_B$(Ag1) = +1.10, $Q_B$(Ag2) = +0.63, and $Q_B$(O) = $-0.89$, compared to the formal values $+3$, $+1$, and $-2$. 

\[ \approx 5.5 \text{ eV} \]

\[ Q_B \approx 0.01 \]

\[ U \approx 5.5 \text{ eV} \]

\[ Q_B \approx 1.45 \]

\[ +2, +4, +3, -2 \]

\[ \approx 0.89 \]

\[ +3, +1, -2 \]
\[ AgNiO_2. \text{ Distorted phase: } Q_B(\text{Ni1}) = +1.15, \quad Q_B(\text{Ni2}) = +1.20, \quad Q_B(\text{Ni3}) = +1.20, \quad Q_B(\text{Ag}) = +0.33, \quad \text{and } Q_B(\text{O}) = -0.84; \quad \text{compare to the formal values } +2.0, +3.5, +3.5, +1.0, \text{ and } -2.0. \]

We have not found the Bader charges in these systems to be useful in identifying and quantifying formal charges. Specifically, the difference in Bader charges for different formal charge states of the transition metal ion are much smaller than the difference in formal charges. The Bader values are consistent with identical ionic charges (3d occupations) but different environments (tails of oxygen 2p orbitals).

As we have surveyed these few systems, we have observed that, if Wannier functions rather than atomic orbitals are used as the physical local orbital, the charge state picture may survive as a useful characterization, at least in most of the cases. The objectivity and generality of this viewpoint remains to be tested, and there are cases such as the RNiO\(_3\) nickelates where the resulting picture seems less than convincing even with Wannier functions in mind. Likely other gray areas will be found as these questions are pursued; fractional charge ordering is especially problematic.

Returning to another underlying motivation—the faithful modeling of ‘charge ordering’ transitions—the basic issues have changed as the microscopic behavior begins to be understood. The initial question we posed was: what are the appropriate terms in a model Hamiltonian to provide correct modeling of the charge, spin and orbital behavior through the transition that leads to ‘charge ordering transitions’? The answer is that this seems not to be the right question. The challenge is different: in a Wannier function basis of 3d orbitals the most transparent local orbital basis, with occupations one or zero, is different on either side of the transition—the environment is instrumental in characterizing the charge state.

The difference can be seen in several ways, but most clearly by the observation that the 3d occupation would differ (by order unity, the difference in occupations) if the orbitals were not considerably different. A physical local orbital description on each side of the transition is simple only if substantially different local orbitals are used on either side. A basis sufficient to enable both representations is the set of all 3d and 2p orbitals (other orbitals are out of the picture); a single simple \( e_g \) or \( t_{2g} \) basis lacks the necessary flexibility. The issue is not what terms are in the Hamiltonian so much as that a good minimal basis on one side of the transition is a poor, insufficient one on the other.
Einstein is said to have stated: ‘a theory (or model) should be as simple as possible, but no simpler’. The requisite basis set, until shown differently, is (1) the set of relevant atomic d states on the open-shell cation (ε₂ or ε₄ when the crystal field splitting is large, for example) and (2) the active 2p atomic orbitals on the ligand (oxygen). These O functions are necessary because the O participation in a Wannier function changes across the transition, and this is a crucial degree of freedom. The on-site repulsion $U$ is relevant due to its role in keeping the d occupation fixed, but perhaps it can be thought of heuristically as a $U \rightarrow \infty$ term as far as the mean-field physics is concerned. Cation–ligand hopping terms and cation–ligand Coulomb energies, both distance dependent, must be important. The on-site energy difference (‘charge transfer’ energy $\varepsilon_q - \varepsilon_p$) of a reference state (the high-symmetry structure) is important in determining the character of the Wannier functions, and Hund’s rule energy is a relevant factor.

Setting up and carrying out fruitful modeling is a task for the future. A fundamental question remains: why does a given charge state—the occupation of a specific type of Wannier orbitals—lead to a reasonably well-defined ionic radius, rather than a continuum depending on the environment—why are ionic radii ‘quantized?’ More broadly stated: why does the conventional charge state picture function so usefully, when 3d (or 4d) charge is not involved—both cation and anion charges are unchanged? An understanding of ‘charge ordering’ transitions may require an answer to this question.

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5 In truth, there is some spread in a given ion’s ‘ionic radius’ in different compounds, but approximate quantization is observed.


