FULL PAPER



Spatial distribution of atomic electronic density for elements 1–54 as coming from a Hartree–Fock treatment within the minimum atomic parameters paradigm

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Peter Reinhardt¹ | Ilya Popov² | Andrei L. Tchougréeff²

¹Laboratoire de Chimie Théorique, Sorbonne Université et CNRS UMR7616, Paris, France

²Frumkin Institute of Physical Chemistry and Electrochemistry, Russian Academy of Sciences, Moscow, Russia

Correspondence

Peter Reinhardt, Laboratoire de Chimie Théorique, Sorbonne Université et CNRS UMR7616, Paris, France. Email: peter.reinhardt@sorbonne-universite.fr

Abstract

The minimum atomic parameters/Moscow-Aachen-Paris (MAP) basis setsreintroduced in the previous paper-are analyzed with respect to spatial features as orbital shape, possible fits to alternative orbital sets (numerical or quasi-numerical orbitals, nodeless Slater orbitals), respect of Kato's condition and radial distribution of energy components. For comparing orbital spaces the Frobenius angle between the orbital subspaces they span is introduced as numerical tool. It is shown that the electronic density of the MAP states is depleted around the nucleus with respect to the other orbital sets. Despite this, the similarity between the respective subspaces in all cases (except a unique case of the Pd atom) as measured by the cosine of the Frobenius angle amounts above 0.96 for all atoms. Deviations from the perfect value of Kato's condition amounts systematically to 0.3 and 0.5 for all elements considered. Integrating one-electron energy contributions from $r = \infty$ to a finite radius, MAP and Bunge orbitals show about the same values, but for the inner region governed by the polynomial oscillations.

KEYWORDS

atoms, Hartree-Fock, minimal parametrization, Slater basis sets

INTRODUCTION. RECALL OF THE MOSCOW-AACHEN-PARIS ORBITALS 1

In the previous paper [1], hereafter referred to as I, we extend the analysis of atomic orbitals constructed via a minimal-parameter procedure, specifying only one single exponent per sub-shell. This procedure-reintroduced in Popov and Tchougréeff [2] for the atoms of the second period-actualizes the original idea dating back to 1930s [3].

In this setting the orthogonality of the atomic functions is obtained by writing the radial function $R_{n\ell}(r)$ as a polynomial of order (n-1) in r times an exponential function $\exp(-\xi_{net})$ and by solving hierarchically the orthonormality conditions for so-constructed functions within a set of common ℓ . Orbital exponents $\xi_{n\ell}$ are chosen to minimize the total energy [4–6] for a given electron configuration, leading to a restricted openshell Hartree-Fock procedure.

As we have shown in paper I, such a setting misses about 3% in total energy with respect to numerical Hartree-Fock orbitals [7, 8], or expansions in the large Slater-type basis sets of Bunge et al. [9], which in their turn give the results at μ -Hartree precision with respect to the previous. Despite this loss in total energy with respect to purely numerical, basis-set-free Hartree-Fock calculations, Moscow-Aachen-Paris (MAP) orbitals have an overlap with their numerical analogues of more than 90%.

As for physical properties, the periodicity of Mendeleev's table is well reproduced by the MAP orbitals in terms of atom sizes. A remarkable feature of the MAP orbitals is that the positions of the nodes in the individual orbitals of a given atom and given angular quantum

2 of 14 WILEY CHEMISTRY

number ℓ almost coincide for different allowed values of the principal quantum number n ($\ell \ge n - 1$) with the coincidence being particularly close for the innermost nodes.

In the present manuscript we investigate further the connections between MAP orbitals and the two other extremes, numerical orbitals on one side and nodeless Slater orbitals on the other, by looking at orbital shapes, mutual fits of parameters, radial integration and Kato's condition as test. Conclusions from these and an outlook to potentially interesting applications close this contribution.

Purely technical details are collected in an Appendix, and a compilation of the data can be found in the accompanying Supporting Information. The underlying MAP orbital exponents are found in the supporting information of paper I [1].

2 | DISCUSSION

All data of this manuscript refer to the electronic ground states of the respective chemical elements, as given by the Madelung–Klechkowski rule [10, 11]. Known exceptions to this rule like Cr, Cu, Pd, and so forth will be discussed in a forthcoming paper, together with a more detailed look at excited states in general.

2.1 | Orbital shapes

Obtained orbitals in the minimal-parameter set may be inspected and compared to available (quasi-)numerical Hartree–Fock orbitals [8, 9]. We observe as general aspect that while the long-range tail is comparable in both orbital sets, the region close to the nucleus manifests significant differences. Specifically, despite the correspondence of the positions of the radial nodes of both the Bunge and MAP functions, the amplitudes of the oscillations of the MAP orbitals coming from the polynomial multiplier are much less pronounced than for Bunge orbitals, especially when comparing valence orbitals, see Figure 1.

Of course, orthonormality imposes the nodal structure of the orbitals, even if the population inside the outermost node, that is, the integral of the squared orbital from zero to the position of this node, does not exceed some percent of the total population of the orbital. This led already Slater to simplify radial functions toward nodeless orbitals [12, 13], and as well for orbitals suited for commonly used pseudopotentials the nodal structure with respect to the replaced core is entirely ignored [14–17]. In the latter case it had been shown that even if Hartree-Fock valence energies are well reproduced, the absence of the inner nodal structure induces significant errors when calculating valence correlation energies. If necessary, one may reduce the number of replaced electrons in order to recover some of the nodal structure (small-core pseudopotentials), or resort to model potentials [18] which are designed to maintain all orbital nodes for the valence orbitals. Since presently we concentrate on the features of the Hartree-Fock based MAP states we keep the discussion of these issues for future work.

Plotting the inner population for MAP and Bunge orbitals as shown in Figure 2 reveals that the spatial distribution of electron density is not the same within the two orbital sets. The difference is relatively small for 2s orbitals, but it becomes quite pronounced for higher *n*. Systematic differences can be seen as well for the series of transition elements (Z = 21-30, 39-48), where the 4s and, correspondingly, the 5s inner population for numerical orbitals—a few percent of the total orbital population—decreases with increasing *Z*, but is hardly detectable for MAP orbitals for these elements.

As we reported in paper I, total energy and atomic radius are affected only weakly by these differences of density distributions, so that overlaps between MAP and Bunge orbitals remain within 95%.

Comparing orbitals in two basis sets one by one does not provide an integral picture. Fundamentally, we should compare functional *subspaces* spanned by the two basis sets. This can be easily done with help of the Frobenius inner product of the projection operators on the respective subspaces (for details see Appendix Equation A1) as a numerical tool. Physically, the cosine of the angle between the subspaces as defined by Equation (A2) gives the probability for a particle (electron) abiding in a state belonging to one subspace to occur in a state of another one.

Figure 3 shows that the so-defined probability remains of the order of 95% throughout the whole range of elements of the periodic table covered here. We observe that Pd represents a striking exception as we optimized the orbital space for a $4d^85s^2$ high-spin state instead of the physical $4d^{10}5s^0$ ground state.

Looking more into detail for the iodine atom (we will use this atom throughout the article as example) we may compare the subspace overlap of Bunge's orbitals with orbital sets generated from a few available basis sets [19]. In Table 1 we give total energies and the Frobenius analysis (cosine of the angle between the subspaces) for the complete subspaces, and broken down for different ℓ . We note that the energy of the MAP orbitals are somewhere between simple STO-nG or double-zeta basis sets. Richer and more flexible basis sets yield better total energies, even if Gaussian basis sets are employed. Measured with the Frobenius analysis, MAP orbitals are on average the least coincident with the target (Bunge) orbitals. Therefore we give in the lower part of Table 1 the overlap matrix elements for each *s* orbital separately, *p* and *d* orbitals following the same trend. We notice that the 1*s* orbital is best represented by the MAP orbital, which is not any more true for the 5*s* orbital. The excellent



FIGURE 1 Illustrating the previous findings at hand of the spatial extension of the *n*s valence orbitals of C, Al, Ni and Ag. Bunge orbitals are represented by solid lines, and MAP orbitals by broken lines. Despite significant differences in the vicinity of the nucleus, the overlap $\int_{0}^{\infty} R_{\text{Bunge}}(r)R_{\text{MAP}}(r)r^{2}dr$ of the corresponding functions is always larger than 90%

representation of the core orbitals by MAP functions (overlaps of 100%) explains the relatively good total energy, and the increasing differences of the polynomial regions with increasing *n* (down to 96% for the 5s orbital) are the reason of or the poorer subspace overlap.

2.2 | Mutual fitting of orbital sets

While numerical Hartree–Fock orbitals can be considered as parameter-free (making abstraction of the radial integration points), we may introduce several different optimization criteria for the parameters defining either MAP or nodeless Slater orbitals.



FIGURE 2 Integration of the radial density from r = 0 up to the outermost node, for Bunge orbitals, and for the corresponding MAP orbitals. Upper panel shows the 2s to 5s, starting at Z = 3, 11, 19, and 37, respectively. Left lower panel displays the *p* series, starting, respectively, at Z = 13, 31 and 49. The 4d orbitals are shown in lower right panel



FIGURE 3 The cosine of the Frobenius angle equation (A2) between the subspaces given by Bunge and MAP orbitals, broken down for different ℓ . The matching of the total orbital space is represented by the continuous line, *s*, *p* and *d* orbital spaces are given by the dash-dotted, dashed and long-dashed lines, respectively. The exception of Pd (Z = 46) is explained in the text

	Total energy	Cos (angle) wrt to Bunge				
Basis	(Hartree)	tot	s	р	d	
STO-3G	-6850.6762	0.9858	0.9852	0.9886	0.9827	
STO-6G	-6885.6062	0.9859	0.9847	0.9892	0.9826	
3-21G	-6887.3052	0.9990	0.9994	0.9988	0.9990	
MAP	-6893.7730	0.9733	0.9794	0.9705	0.9735	
6-311G	-6904.0420	0.9608	0.9971	0.9968	0.8995	
Jorge DZP	-6911.9867	0.9906	0.9994	0.9973	0.9783	
6-311G**	-6916.7496	0.9995	1.0000	0.9999	0.9987	
Jorge QZP	-6917.5145	0.9998	1.0000	0.9999	0.9996	
Bunge	-6917.9809					
	Overlap matrix	elements				
Basis	<u>1</u> 5	2s	3s	4s	5s	
STO-3G	0.9998	0.9979	0.9952	0.9923	0.9754	
STO-6G	1.0000	0.9967	0.9930	0.9912	0.9751	
3-21G	0.9995	0.9997	0.9997	0.9999	0.9998	
MAP	1.0000	0.9985	0.9895	0.9777	0.9589	
6-311G	1.0000	1.0000	0.9999	0.9980	0.9933	
Jorge DZP	1.0000	0.9999	1.0000	0.9995	0.9984	
6-311G**	1.0000	1.0000	1.0000	1.0000	1.0000	
Jorge QZP	1.0000	0.9999	1.0000	1.0000	1.0000	

Note: The upper part of the table gives the different subspaces (total, *s*, *p* and *d* orbitals). Orbital sets are ordered wrt to total energy. In the lower part of the table we give the individual overlap-matrix elements $\langle \phi_i^{\text{basis}} | \phi_i^{\text{Bunge}} \rangle$ for the five different *s* orbitals.

2.2.1 | Orbital exponent optimization with nodes fixed at the positions of the nodes given by numerical orbitals

Having observed that the orbital shapes in the core region are significantly different between the MAP and Bunge/numerical orbitals, despite the relative smallness of the total-energy differences and large overlaps, we may have a closer look to the nodes of the orbitals. As we could see for the iodine 5s orbital as example (see figure 5 of paper I), the nodes are nearly coinciding, although not identical between MAP and Bunge orbitals.

The numerical orbital has, due to the orthogonality constraints, as many nodes as the corresponding MAP orbital. We may try to fit thus the MAP expression to this orbital, going through exactly the same nodes. Before doing so, we divide numerically the radial function of numerical orbitals by the corresponding polynomial factor. As shown in Figure 4, the ideally linear relations (in a logarithmic plot) are well visible for the inner shells, and as well the shell structure is clearly reproduced.

Fitting a MAP orbital to the corresponding numerical Hartree–Fock or Bunge orbital while keeping the node positions fixed at those of the target orbital, comes then down to a linear regression with the slope (i.e., the MAP exponent) as only free parameter. The intercept is fixed through the normalization condition.

Optimizing the overlap of MAP states in this way gives for instance for the iodine 5s orbital an exponent of 3.18631 with overlap of 0.964, almost identical with one found via the energy optimization in the MAP setting: 3.1897. A more detailed compilation of the results of the described procedure applied to the whole set of iodine orbitals is given in Table 2.

Taking the so-obtained, slightly non-orthogonal orbitals—we fit the MAP exponents individually to the corresponding Bunge orbitals—as starting point for our MAP energy optimization procedure, we obtain again the previously generated, energy-minimizing exponents.

2.2.2 | Fit MAP to Bunge by maximizing the Frobenius product between the two orbitals sets

As mentioned in Section 2.1, the numerical measure of the coincidence between the orbital subspaces spanned by different basis sets is given by the Frobenius angle equation (A2) between them. The cosine of this angle is the probability of finding an electron in a subspace spanned by one

TABLE 1Frobenius analysis fordifferent standard basis sets for an iodineatom, with respect to Bunge orbitals

WILEY 5 of 14





FIGURE 4 For iodine we show the numerical Hartree–Fock orbitals—as obtained by the Froese–Fischer program [8]—divided by the polynomial fixed by the orbital nodes, in a logarithmic scale. The 1*s* orbital is a pure exponential function; however the deviations from the ideal relations are more and more pronounced for increasing *n*

Orb.	MAP exp. var. HF	MAP/Slater overlap	MAP exp. fitted to Bunge	Max. overlap in %
1 <i>s</i>	52.686	100.	52.3059	99.9993
2 <i>s</i>	25.249	87.8	24.2894	99.9565
3s	14.883	72.1	13.8225	99.5837
4 <i>s</i>	8.1798	71.1	7.65756	98.3884
5s	3.1897	86.4	3.18631	96.4233
2p	24.632	100.	23.7707	99.9710
Зр	14.276	81.8	13.3155	99.5463
4p	7.5596	77.5	7.05405	98.1482
5p	2.7051	72.0	2.58773	95.9579
3d	12.988	100.	12.1820	99.7152
4 <i>d</i>	6.1826	89.8	5.72662	97.5304

TABLE 2 MAP exponents for an iodine atom

Note: The first column gives the previously obtained MAP exponents (see supporting information of paper I for full-precision data), second column gives the overlap (in per cent) of the MAP function and a corresponding nodeless Slater orbital $r^{(n-1)}e^{-\xi r}$. Third and fourth column show the data for the exponents obtained by maximizing the overlap of the same $n\ell$, respecting norm and positions of the nodes. Total energies are (in Hartree) -6917.981, -6893.773, -6881.287 for Bunge, variational MAP, and the present fit, respectively. For calculating a total energy from the fitted exponents, orbitals were re-orthogonalized via the MAP construction. This shifts the node positions slightly, as Bunge nodes and orthogonality define too many constraints to be satisfied simultaneously within the minimal parameter space.

basis set under the condition that it occurs in the subspace spanned by another one. We may thus maximize this quantity by fitting all exponents of the mutually orthogonal MAP orbitals to Bunge orbitals, without the exact node-matching.

Traditionally, (see, e.g., References [5, 20–23]) the quality of a fit is controlled by the achieved total energy as based on the variation principle. This approach is actually not needed. First, it requires the calculation of the energy which may be tedious. Second, in the strict sense, it produces the measure of the closeness of the specific (lowest energy) state approximations achievable in the subspaces rather than of the subspaces themselves. In the atomic context it means that in the energy-variation principle setting the states, say, in the valence subshell of an atom, are treated differently depending on the amount of electron population of the latter. Thus, the Frobenius angle or its cosine gives a more uniform quantification of the similarity of the subspaces to be compared.

Figure 5 gives an account on the difference of the two optimization criteria, and the corresponding exponents can be found in the Supporting Information. In contrast to energy optimization we face now 1s exponents which are larger than the atomic number *Z*, especially for the heavier elements, for example, for iodine we arrive at 53.2598. One explanation may be that Bunge's basis sets contain as well exponents larger than the atomic number, with non-negligible weight in the 1s orbitals. However, the same results are obtained when maximizing the overlap with respect

to basis-set free numerical orbitals—we cannot impute this apparently unphysical result to numerical artifacts, but to the optimization criterion which gives less relative weight to the core orbitals when maximizing the overlaps globally.

2.2.3 | Fit of Z^{*} of the Slater model to MAP and Bunge orbitals

The Frobenius scalar product allows to quantify the difference between different orbital sets. Another measure may be to fit a well-defined third orbital class to either set, for instance Slater's nodeless radial functions, by maximizing the overlap between the orbital sets.

Again, we have just one single parameter per subshell, as all others are fixed.

The overlap of a MAP or Bunge orbital with a primitive Slater orbital may be easily calculated as a function of Z^* with the help of, for example, Mathematica [24], and incorporated in an optimization procedure.

For the iodine atom as example we find the best Z^{*} yielding the maximal overlaps as given in Table 3. The complete sets of orbital exponents can be found in the Supporting Information.

We see immediately that overlaps between MAP and Bunge states are always better than 0.95, and the obtained Z^* are close. However, the resulting 2s exponent is considerably smaller than the 2p exponent, and the same holds for the 3p exponent. For these two orbitals the relatively large oscillations of the polynomial render the best-overlap nodeless Slater functions shifted with respect to the main lobes. A larger exponent would fit the outer region better, but gives a smaller overlap in the inner region. For the higher-*n* orbitals this is less important as the polynomial oscillations are weaker, and thus the common decay at larger distances is more significant for the best overlap.

Figure 6 gives a global picture of the results of the two fitting procedures, in terms of analogues of effective screenings $S_{n\ell}$ of the nucleus

$$\xi_{n\ell} = \frac{Z_{n\ell}^*}{n^*} = \frac{Z - S_{n\ell}}{n^*}$$

The *sp* degeneracy of Slater's model is lifted, and the screenings always augment with augmenting *Z* (exceptions being the nodeless 1s, 2*p* and 3*d* shells). The order of magnitude (i.e., atomic radii) is the same for all three orbital sets. Having a closer look to the 3*d* orbital, we may notice a qualitatively different evolution of MAP and Bunge orbitals, the MAP orbital being more compact than the corresponding one from Bunge's set.

2.3 | Kato's condition

As commonly known, the exact electron density must have a cusp at the nuclear position, respecting Kato's condition: [25]

$$\frac{1}{2} \frac{\left|\vec{\nabla}\rho(\vec{r})\right|_{\text{sph.av.}}}{\rho(\vec{r})} \bigg|_{r=0} = Z.$$
(1)



FIGURE 5 Cosine of the Frobenius angle wrt to Bunge orbitals for the energy-optimized MAP orbitals (solid line) and the reoptimized orbitals to minimize this angle. The spike for Pd (Z = 46) disappears, as we do not optimize the total energy for a given state, but for a given set of orbitals

8 of 14 WILEY CHEMISTRY

TABLE 3 Slater's effective nuclear charges Z^* and maximized overlaps and corresponding Z^* for MAP/Bunge orbitals and Slater orbitals $r^{(n^*-1)}\exp(\frac{Z^*}{n^*}r)$, for the iodine atom, Z = 53

		МАР		Bunge		
Orb.	Slater's screened Z [*]	Max. overlap	Z [*]	Max. Overlap	Z	S_{MAP/Bunge}
1s	52.69	1.000000	52.6862	0.999993	52.3059	0.999973
2 <i>s</i>	48.85	0.932635	38.8031	0.938749	37.3943	0.998507
3s	41.75	0.882507	30.8542	0.895175	28.4649	0.989460
4s	25.25	0.896528	19.6500	0.893065	17.8801	0.977660
5s	7.60	0.947849	8.13347	0.942856	7.97763	0.958939
2p	48.85	1.000000	49.2630	0.999710	47.5415	0.998949
Зр	41.75	0.910455	32.2099	0.919719	29.7338	0.990418
4p	25.25	0.914354	18.9800	0.911075	17.2700	0.976573
5p	7.60	0.958761	7.13512	0.959876	6.82707	0.954011
3d	31.85	1.000000	38.9644	0.997152	36.5458	0.993981
4d	13.85	0.953058	17.2792	0.948578	15.8040	0.972777

Note: We give in the last column also the overlap between the MAP and Bunge orbital for sake of completeness.



FIGURE 6 Scaled screenings of the nucleus $\left(-\frac{S_{nc}}{n^*} = \frac{Z_{nc}^* - Z}{n^*}\right)$, for the Slater model (left), Slater functions fitted to MAP orbitals (middle) and to Bunge orbitals (right), as functions of the atomic number Z. The constant screenings and the sp degeneracy of Slater's model are not reproduced, but for the nodeless 1s, 2p and 3d orbitals. For MAP and Bunge orbitals we give as well (dashed lines) the screenings scaled with the full quantum number *n* instead of Slater's *n*^{*}. Shells can be identified in the middle and right panel from the starting points of the lines. The spike in the right panel is Pd (Z = 46)

Taking numerical Hartree–Fock orbitals or Bunge's basis sets we see that this condition is fulfilled already at the Hartree–Fock level [7], and even individually for each atomic *s* orbital [26]—the only ones with a non-vanishing electron density at the origin.

Despite the sizable absolute deviation of the gradient-to-density ratio from the ideal value of *Z* it does not show any significant or characteristic variation with *Z*, ranging from 0.3 for a helium atom to around 0.5 for the heavier elements. The relative deviation (divided by *Z*) fades out with increase of *Z*. Electron correlation as the missing ingredient in the Hartree–Fock approach can be ruled out being responsible for this deviation as the electron density and orbitals within Bunge's basis sets reproduce the nuclear cusp correctly.

We find that the 1s exponent already governs the value of the gradient at the origin. The other s orbitals change this quantity only very little, as they add in a similar way to the numerator and the denominator of the expression. For instance for the Sulfur atom (Z = 16) we find as

individual $\left|\vec{\nabla}\rho(\vec{r})\right|/\rho(\vec{r})\right|_{r=0}$ values of 15.6886, 14.1127 and 11.1905 for 1s, 2s, and 3s, respectively, leading as a sum to 15.6052. Using Bunge's basis sets the corresponding data are 15.9999, 15.9919 and 15.9976 with a sum of 15.9993, very close to the expected result.

The overall results, as cumulative sums in numerator and denominator of the reduced density gradient, are depicted in Figure 7; the smooth curves allow an estimation of the numerical noise of our multi-parameter optimization procedure.

We see as well that the 4s and the 5s orbitals do not add any change to the cumulative sums. This is due to the very low electron density at the origin, for instance for the I atom in the MAP setting we have densities of 7499, 757.5, 99.83, 3.100, and 0.001045 (in atomic units) for 1s to 5s, respectively. For orbitals expanded in the Bunge basis set, the figures read 7391, 779.5, 154.7, 31.45 and 3.639, reflecting the same findings as for the populations inside the polynomial region.

2.4 | Radial integration of one-electron quantities

Kato's condition "measures" the orbitals in the vicinity of the nucleus. For looking at the opposite side, we may integrate energy contributions [7] from outside toward the nucleus, like kinetic energy or the electron-nucleus attraction. In this way we see, in which region the difference in total energy between the MAP and the Bunge orbital sets are located. Figure 8 shows these contributions for the orbitals already displayed in Figure 1. If for the carbon 2s orbital the difference is hardly visible, we see for the Ni 4s and Ag 5s orbital that the kinetic-energy contributions to the total energy are not systematically underestimated by the MAP orbitals,

Radii for the shown elements (C, Al, Ni, Ag) are 1.94025, 3.34261, 3.20627 and 3.63247 Bohr, respectively, and nodes of Bunge orbitals are situated at 0.376 a.u. (C, 2s), 0.797 and 0.162 (Al, 3s), at 0.0742 (Ni 2s), 0.304 and 0.0718 (Ni, 3s), 0.936, 0.296 and 0.0716 (Ni, 4s), and, finally, at 1.149, 0.439, 0.162, and 0.0417 a.u. (Ag, 5s). We marked these in the plot.

In the region outside the atomic radius the two orbital sets are close, and significant deviations for the radial integration occur only within the nodal region, too close to the nucleus with respect to the chemically important valence region. We can thus expect that atoms described with MAP orbitals will produce chemically relevant data, without the need for long basis set expansions. The density missing in the core region is distributed in a wide range in the outer orbital lobes, rendering them only insignificantly different to numerical or high-precision orbitals, be it in Gaussian or Slater expansions.

2.5 | Relation to the general basis-construction problems

Depending on the application, a typical ab initio Gaussian basis set employs 10–200 GTO functions per atom, optimized with some criterion but without whatever individual meaning [26]. Yet at pretty early stage there were attempts to reduce the number of independent parameters in soconstructed sets by subjecting the exponents of the *Gaussians* to some parametric form. After the STO-nG series [27], this gave rise to welltempered and even-tempered basis sets [20–23], well known in the literature. Remarkably, the functions of these sets have been obtained by



9 of 14

M→WILEY



FIGURE 8 The integral from R to ∞ of the kinetic energy (positive) and e-n one-electron contribution to the potential energy (negative), for the six orbitals of Figure 1. We mark as vertical lines the positions of the orbital nodes, and for the valence orbitals with a broken vertical line the atomic radius

minimizing the total energy in the Hartree–Fock approximation by which basis sets eventually spanning the same subspace as the Bunge basis sets, the latter very close to numerical Hartree–Fock orbitals obtained on radial grids [8]. The parameters used for the construction of even- or well-tempered exponent sequences, universal [20, 22] or atom-specific [23], do not have any meaning by themselves—they are merely numbers.

The present MAP paradigm provides a completely different approach to the selection of the characteristics of the spatial decay of the atomic states: for each (sub)shell a single parameter provides a definite physical sense—a great intellectual and technical advantage of the proposed approach. So far implemented for the orbitals occupied in the ground state as treated in the Hartree–Fock setting, the MAP approach may be extended to excited (unoccupied in the ground state) atomic orbitals to formation of hierarchical basis sets as prescribed in Blum et al. [28]. Indeed, adding further and further subshells entering with corresponding exponents with assured orthogonality eventually produces the required hierarchy of approximations.

This still needs to be controlled for convergence properties, and tested for reasonable sources of the additional exponents for the subshells missing in the ground state. We will address this issue in a future work.

UANTUM WILEY 11 of 14

Although, maybe not well suitable for describing subtle physical effects depending on the details of the function behavior in the vicinity of the nucleus like magnetic properties and/or electric (hyper)polarizabilities, the very reasonable exponential decay at larger distances from the nuclei, important for bond formations and chemical reactivity, makes the MAP states a good alternative for studying chemistry. The paradigm may be extended as well to ionic states, and thus charge-transfer and ionic compounds. Instead of density plots expanded in a large number of Gaussian basis functions, a few orbital exponents may be sufficient to characterize different chemical situations.

3 | CONCLUSIONS

By the present work, we continue our studies on minimally parameterized atomic states extending them either in terms of rows covered of the Mendeleev periodic table. We concentrate particularly on the spatial behavior of the MAP states as compared to that of traditional Slater orbitals and quasi-numerical Bunge orbitals. It turns out that the MAP states, although, showing much weaker oscillation amplitudes at the shorter electron-nucleus separations, manifest very large (typically more than 0.95) overlaps with the Bunge states on account of the almost coinciding positions of the radial nodes of the either series of orbitals. At the same time the similarity of the MAP states with the Slater orbitals allows to qualify the former as minimally orthogonalized Slater orbitals. In this quality the MAP states are perspective for developing new semi-empirical methods which habitually use the Slater basis sets which due to lack of the radial nodes cannot assure correct relative positioning of the states of the same *n* but different ℓ on the energy scale. These and other issues and perspectives will be addressed in future work.

Additionally, we propose a numerical tool for comparing the relative quality of the basis sets through the Frobenius inner product of the operator (matrices) projecting to the subspaces spanned by the basis sets to be compared.

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AUTHOR CONTRIBUTIONS

Peter Reinhardt: Software; data acquisition; writing the manuscript. **Ilya Popov:** Data curation; formal analysis; resources; software; validation. Andrei Tchougreeff: Conceptualization; formal analysis; methodology; project administration; supervision; writing-original draft.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

ORCID

Peter Reinhardt D https://orcid.org/0000-0001-8644-7468

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12 of 14 WILEY-CHEMISTRY

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SUPPORTING INFORMATION

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APPENDIX A: Technical details

The Frobenius inner product

Given a basis of the Hilbert space of L^2 integrable functions we may define the projectors on the subspaces given by two (finite) orthonormal sets of atomic orbitals, as $\hat{M} = \sum_{\mu=1}^{m} |\mu\rangle \langle \mu|$ and $\hat{B} = \sum_{\beta=1}^{b} |\beta\rangle \langle \beta|$, respectively. The trace of the product of the two matrices tr($M^{\dagger}B$) defines a scalar product (inner Frobenius product) in the space of matrices for which the usual matrix multiplication is possible, making the latter an Euclidean vector space. It has all properties of a scalar product—bilinearity and positive definiteness for M = C, but for M being the zero matrix. Applying it to the square (projector) matrices $(M)_{\lambda\kappa} = \langle \lambda | \hat{M} | \kappa \rangle$ and B yields:

$$\operatorname{tr}(M^{\dagger}B) = \sum_{\kappa\lambda} \sum_{\mu} \sum_{\beta} \langle \kappa | \mu \rangle \langle \mu | \lambda \rangle \langle \lambda | \beta \rangle \langle \beta | \kappa \rangle$$

$$= \sum_{\mu} \sum_{\beta} \left\langle \beta \sum_{\kappa} | \kappa \rangle \langle \kappa | \mu \rangle \right\rangle \left\langle \mu \sum_{\lambda} | \lambda \rangle \langle \lambda | \beta \rangle$$

$$= \sum_{\mu} \sum_{\beta} |\langle \beta | \mu \rangle|^{2},$$
(A1)

where $|\lambda\rangle$ and $|\kappa\rangle$ run over a complete orthonormal basis in L². That matrix elements are squared in the above expression does not destroy the bilinearity of the scalar product, as the underlying space is not the orbital space, but the space of the matrices acting in the Hilbert space.

A norm is derived as usual as the square root of the Frobenius inner product of a matrix with itself: $|C| = \sqrt{tr(C^{\dagger}C)}$ and is known as the Frobenius matrix norm. The angle between two orbital subspaces is defined via its cosine as:

$$\cos\varphi_{\rm F} = \frac{\operatorname{tr}(M^{\dagger}B)}{|M||B|} = \frac{\sum_{\mu=1}^{m} \sum_{\beta=1}^{b} |\langle\beta|\mu\rangle|^2}{\sqrt{m}\sqrt{b}}.$$
 (A2)

As the involved orbital sets (MAP and Bunge in the present case) are orthonormal, the inner product of either projector with itself is just the dimensionality of the subspaces *m* or *b* (number of elements in the respective orbital set, taking into account the multiplicity of $2\ell + 1$ for a subshell with the azimuthal quantum number ℓ) and the Frobenius norm is the square root of it. The expression Equation (A2) is always non-negative, and falls between 0 and 1. It can be connected to the notion of a probability.

Integration of energy contributions

For integrating one-electron contributions to the total energy it is sufficient to look at the radial functions. For the kinetic energy the radial part of the Laplace operator reads

$$\frac{d^2}{dr^2} + \frac{2}{r}\frac{d}{dr} - \frac{\ell(\ell+1)}{r^2}$$

which yields for a pair of primitive, unnormalized Slater functions for a given ℓ

$$r^{(m-1)}e^{-\beta r}\Delta_{r}r^{(n-1)}e^{-\alpha r} = r^{(m-1)}e^{-\beta r}\left(\frac{d^{2}}{dr^{2}} + \frac{2}{r}\frac{d}{dr} - \frac{\ell(\ell+1)}{r^{2}}\right)r^{(n-1)}e^{-\alpha r}$$

$$= e^{-(\alpha+\beta)r}r^{(m+n-2)}\left(\frac{n(n-1)-\ell(\ell+1)}{r^{2}} + \alpha^{2} - \frac{2}{r}n\alpha\right)$$
(A3)

For obtaining the kinetic energy density we have to multiply this with -1/2. We may notice that for instance for $\ell = n - 1$ the density of the kinetic energy becomes negative for $r > (2n)/\alpha$.

Integrating a product of unnormalized Slater functions from 0 to R gives

$$\int_{0}^{R} r^{(m-1)} e^{-\beta r} r^{(n-1)} e^{-\alpha r} r^{2} dr = \int_{0}^{R} r^{(m+n)} e^{-(\alpha+\beta)r} dr =$$

$$= \left(\frac{1}{\alpha+\beta}\right)^{m+n+1} \left((m+n)! - \underbrace{\Gamma(m+n+1,R(\alpha+\beta))}_{-0 \text{ for } R \to \infty} \right)$$
(A4)

For the potential we find thus

$$\int_{0}^{R} r^{(m+n-1)} e^{-(\alpha+\beta)r} dr = \left(\frac{1}{\alpha+\beta}\right)^{m+n} \left((m+n-1)! - \underbrace{\Gamma(m+n,R(\alpha+\beta))}_{\rightarrow 0 \text{ for } R\rightarrow\infty} \right)$$
(A5)

 $\Gamma(a,z) = \int_{z}^{\infty} t^{(a-1)} e^{-t} dt$ is the incomplete Γ its value to high precision. We may recall that the normalized Slater function $S_{n}^{\alpha}(r)$ is

$$S_n^{\alpha}(r) = \frac{(2\alpha)^{3/2}}{\sqrt{\Gamma(2n+1)}} (2\alpha r)^{n-1} e^{-\alpha r}$$
(A6)

For the radial integration of the kinetic energy the expressions become slightly more complicated:

$$\begin{split} & -\frac{1}{2} \int_{0}^{R} S_{m}^{\beta}(r) \left(\frac{d^{2}}{dr^{2}} + \frac{2}{r} \frac{d}{dr} - \frac{\ell(\ell+1)}{r^{2}} \right) S_{n}^{\alpha}(r) r^{2} dr \\ & = \frac{2^{m-1} \beta^{m+1/2}}{\sqrt{m \Gamma(2m) \Gamma(2m)}} \times \left(-\frac{2^{n} n(n-1) \alpha^{n+3/2} (m+n-2)!}{(\alpha+\beta)^{m+n}} + \frac{\ell(\ell+1)}{\beta^{m}} \sqrt{n \Gamma(2n)} \left(\Gamma(m) - \underbrace{\Gamma(m, R\beta)}_{-0 \text{ for } R \to \infty} \right) - \frac{2^{n} e^{-R(\alpha+\beta)} \alpha^{n+1/2} R^{m+n-1} (n(n-1)(\alpha+\beta) - (m+n-1)R\alpha^{2})}{(m+n-1)(\alpha+\beta)} + \frac{2^{n} \alpha^{n+1/2}}{(m+n-1)(\alpha+\beta)^{m+n+1}} \times \\ & \left\{ \left((m-m^{2}) \alpha^{2} + 2mn\beta + n(n-1)(\alpha^{2} + \alpha\beta - \beta^{2}) \right) (m+n-1)! + \frac{(m(m-1)\alpha^{2} - 2mn\alpha\beta + n(n-1)\beta^{2}) \Gamma(m+n, R(\alpha+\beta))}{-0 \text{ for } R \to \infty} \right\} \end{split}$$

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$$-\frac{1}{2}\int_{0}^{\infty} r^{(m-1)} e^{-\beta r} \left(\frac{d^{2}}{dr^{2}} + \frac{2}{r}\frac{d}{dr} - \frac{\ell(\ell+1)}{r^{2}}\right) r^{(n-1)} e^{-\alpha r} r^{2} dr$$

$$= \frac{2^{m-1} \beta^{m+1/2}}{\sqrt{mn\Gamma(2m)\Gamma(2n)}} \times \left(-\frac{2^{n} n(n-1) \alpha^{n+3/2} (m+n-2)!}{(\alpha+\beta)^{m+n}} + \right)$$
(A8)

$$\frac{\ell(\ell+1)}{\beta^{m}}\sqrt{n\Gamma(2n)}\Gamma(m) + \frac{2^{n}\alpha^{n+1/2}}{(\alpha+\beta)^{m+n+1}} \times \left(\left(m-m^{2}\right)\alpha^{2} + 2mn\beta + n(n-1)\left(\alpha^{2} + \alpha\beta - \beta^{2}\right)\right)(m+n-2)!\right)$$

$$=\frac{2^{m-1}\sqrt{\beta}(m-1)!}{\sqrt{m(2m-1)!}}\ell(\ell+1) - \frac{2^{m+n-1}\alpha^{n+1/2}\beta^{m+1/2}}{\sqrt{mn\Gamma(2m)\Gamma(2n)}(\alpha+\beta)^{m+n+1}}(m+n-2)!\times$$

$$\left((m-m^{2})\alpha^{2} + 2mn\beta + n(n-1)(\alpha^{2}+\alpha\beta-\beta^{2}) - n(n-1)\alpha(\alpha+\beta)\right)$$
(A9)

$$=\frac{2^{m-1}\sqrt{\beta}(m-1)!}{\sqrt{m(2m-1)!}}\ell(\ell+1) - \frac{2^{m+n-1}\alpha^{n+1/2}\beta^{m+1/2}(m(m-1)\alpha^2 - 2mn\beta + n(n-1)\beta^2)}{\sqrt{mn\Gamma(2m)\Gamma(2n)}(\alpha+\beta)^{m+n+1}}(m+n-2)!$$
(A10)

Mathematica gives for the latter the same two terms.