

Assessment of Tuning Methods for Enforcing Approximate Energy Linearity in Range-Separated Hybrid Functionals

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Abstract

A range of tuning methods, for enforcing approximate energy linearity through a system-by-system optimisation of a range-separated hybrid functional, are assessed. For a series of atoms, the accuracy of the frontier orbital energies, ionisation potentials, electron affinities, and orbital energy gaps is quantified and particular attention is paid to the extent to which approximate energy linearity is actually achieved. The tuning methods can yield significantly improved orbital energies and orbital energy gaps, compared to those from conventional functionals. For systems with integer M electrons, optimal results are obtained using a tuning norm based on the highest occupied orbital energy of the M and $M + 1$ electron systems, with deviations of just 0.1 – 0.2 eV in these quantities, compared to exact values. However, detailed examination for the carbon atom illustrates a subtle cancellation between errors arising from

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non-linearity and errors in the computed ionisation potentials and electron affinities used in the tuning.

1 Introduction

A significant problem in Kohn–Sham density functional theory¹ (DFT) is the delocalisation error,^{2–4} present in commonly-used exchange–correlation functionals. This problem underlies many of the well-known failures of approximate DFT, including the underestimation of quantities such as reaction barrier heights,⁵ band gaps,⁶ energies of dissociating molecular ions,^{2,3,7,8} and charge–transfer excitation energies.⁹

Delocalisation error can be illustrated by considering the variation in the total electronic energy, E , as a function of electron number, N . Perdew et al.¹⁰ showed that the exact E varies linearly with fractional N , with discontinuities in the gradient $\frac{\partial E}{\partial N}$ at integer N . The overall E vs. N behaviour is therefore a piecewise function comprising a series of linear segments between the integers. Functionals that suffer from delocalisation error do not exhibit this piecewise linear behaviour—instead they give a convex deviation from linearity, such that the energy at fractional N is too low. As a result, these functionals favour locally fractional charge, over-delocalising systems and underestimating their energy. Such deviation from linearity is also known as many-electron self-interaction error.^{11–16}

This erroneous non-linear behaviour has important implications for the Kohn–Sham orbital energies and their relationship to ionisation potentials and electron affinities. Let M be an integer and $0 < f < 1$. The piecewise linearity of the exact E vs. N curve implies that for a system with $N = M - f$ electrons, the exact $\frac{\partial E}{\partial N}$ is equal to the negative of the exact vertical ionisation potential of the M -electron system, denoted I_0^M , independent of the value of f . Similarly, for a system with $N = M + f$ electrons, $\frac{\partial E}{\partial N}$ equals the negative of the exact vertical electron affinity of the M -electron system, denoted A_0^M , again independent of f . From Janak’s theorem,¹⁷ $\frac{\partial E}{\partial N}$ equals the energy of the

Kohn–Sham orbital whose occupation is varying. It follows that

$$\epsilon_M(M - f) = -I_0^M, \tag{1}$$

$$\epsilon_{M+1}(M + f) = -A_0^M, \tag{2}$$

where $\epsilon_M(M - f)$ is the energy of the M th orbital of the $M - f$ electron system and $\epsilon_{M+1}(M + f)$ is the energy of the $(M + 1)$ th orbital of the $M + f$ electron system. In the limit $f \rightarrow 0$, eqs. 1 and 2 can be identified as the exact Koopmans relationships

$$\epsilon_{\text{H}}^{M,-} = -I_0^M, \tag{3}$$

$$\epsilon_{\text{L}}^{M,+} = -A_0^M, \tag{4}$$

where $\epsilon_{\text{H}}^{M,-}$ and $\epsilon_{\text{L}}^{M,+}$ are the exact highest-occupied (HOMO) and lowest-unoccupied (LUMO) molecular orbital energies of the M -electron system, determined on the $f \rightarrow 0$ electron-deficient and electron-abundant sides of the integer, respectively. The use of the $+/-$ superscript to denote the side of the integer is vital, because the exact exchange–correlation potential jumps discontinuously as the integer is crossed, meaning a given orbital energy also jumps by the same amount. In practical calculations, using approximate exchange–correlation functionals within the usual generalised Kohn–Sham approach,¹⁸ $\frac{\partial E}{\partial N}$ is again equal to the orbital energy⁶ and so the value of $\frac{\partial E}{\partial N}$ on the $f \rightarrow 0$ electron-deficient and electron-abundant sides of the integer M equals the HOMO energy, ϵ_{H}^M , and the LUMO energy, ϵ_{L}^M , of the M -electron system, respectively. However, the incorrect E vs. N curvature associated with the delocalisation error means that $\epsilon_{\text{H}}^M \gg -I_0^M$ and $\epsilon_{\text{L}}^M \ll -A_0^M$ (Refs. 6,19).

Several approaches can be used to impose near-linear E vs. N behaviour. Vydrov et al.¹² demonstrated that many electron self-interaction error was significantly reduced by applying a self-interaction correction;²⁰ see also Refs. 21–23. The MCY3 and rCAM-B3LYP functionals¹⁴ were specifically designed to achieve near-linear behaviour and they have shown some success.²⁴ Zheng et al.²⁵ proposed a non-empirical scaling correction to largely restore linearity, which was

later extended²⁶ to properly account for orbital relaxation effects. Very recently, Kraisler and Kronik²⁷ demonstrated that the exact Koopmans ionisation relationship could be largely restored using an ensemble treatment. A more pragmatic approach is to simply tune the proportion of exact orbital exchange in global^{28–30} or range-separated^{31–38} hybrid functionals on a system-by-system basis, in order to approximately recover Koopmans or other conditions, which are necessary (but not sufficient) for linearity. The success of this approach reflects the fact that Hartree–Fock theory typically yields concave E vs. N curves, and so the inclusion of exact orbital exchange tends to cancel the incorrect convex behaviour. Karolewski et al.³⁹ recently highlighted a number of caveats for such an approach, most notably the violation of size-consistency. The aim of the present study is to provide a systematic investigation of a wide range of tuning methods, quantifying the accuracy of Koopmans relationships, ionisation potentials, electron affinities, and orbital energy gaps. Particular attention is paid to the degree of linearity of the resulting E vs. N curves. Results and Conclusions are presented in Sections 2 and 3, respectively.

2 Results

2.1 Choice of Tuning Methods

Figure 1 illustrates the key quantities in the context of an E vs. N diagram. The dashed blue curve represents the convex behaviour of a typical approximate exchange–correlation functional within the generalised Kohn–Sham approach. The slopes of this curve on the limiting electron-deficient side of the M - and $(M + 1)$ -electron integer systems are the HOMO energies of those systems, ϵ_{H}^M and $\epsilon_{\text{H}}^{M+1}$, respectively. The slopes on the limiting electron-abundant side of the $(M - 1)$ - and M -electron systems are the LUMO energies of those systems, $\epsilon_{\text{L}}^{M-1}$ and ϵ_{L}^M , respectively. Also indicated are the vertical ionisation potentials, I^M and I^{M+1} , and electron affinities, A^{M-1} and A^M , of the integer systems, determined as the differences between the energies of the integer systems, calculated using the same approximate functional. The solid black curve indicates a linear interpolation between the integer energies, which will more closely resemble the shape of the exact

curve because approximate functionals tend to perform more accurately at integer electron-number. For brevity, we denote the E vs. N curve from $M - 1 \rightarrow M$ as the left-hand side (LHS) and the curve from $M \rightarrow M + 1$ as the right-hand side (RHS).

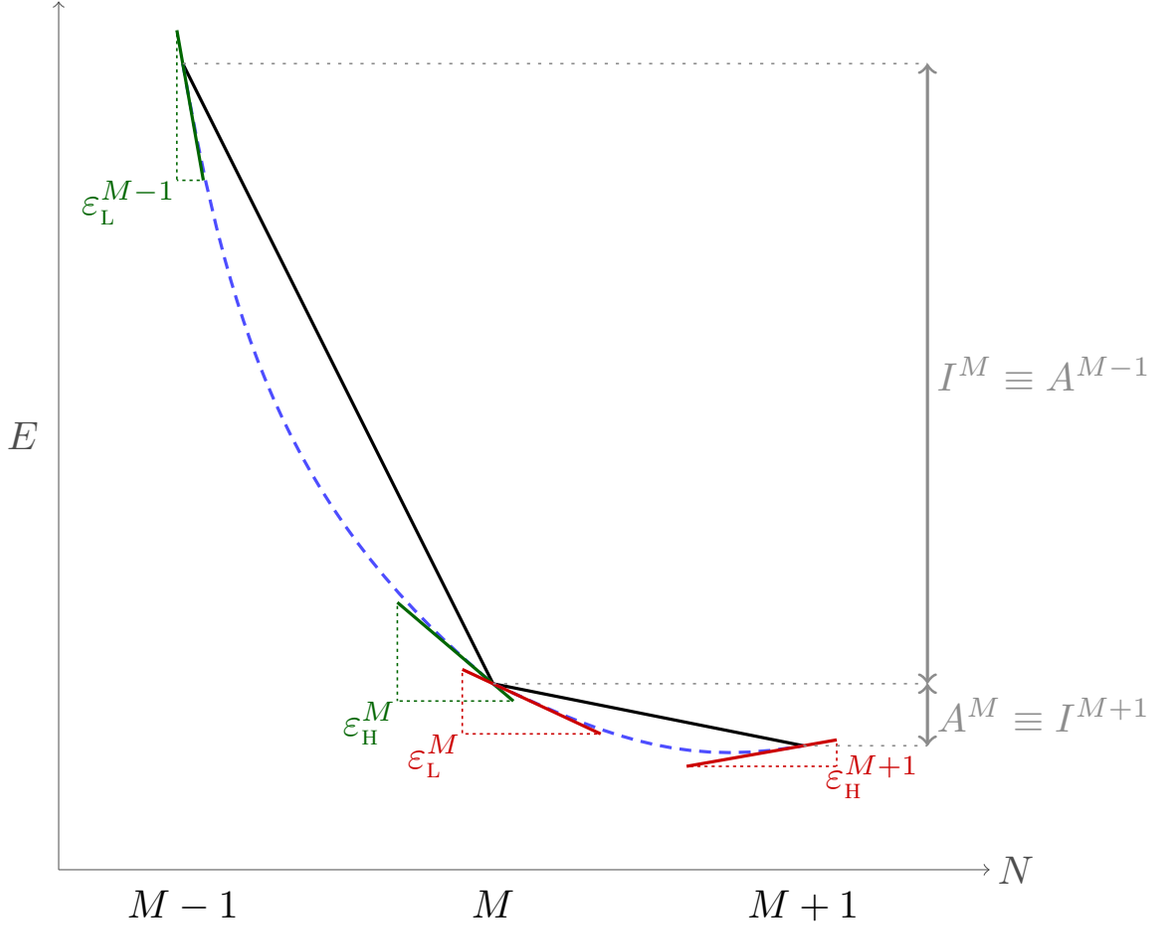


Figure 1: Schematic representation of the quantities involved in the tuning methods.

In the context of Figure 1, the aim of a tuning method is to make the non-linear blue curve more closely resemble the piecewise linear black one. If this could be achieved on the LHS then the following condition would hold,

$$\epsilon_H^M = \epsilon_L^{M-1} = -I^M = -A^{M-1}, \quad (5)$$

which suggests three possible tuning criteria:

$$\epsilon_{\text{H}}^M = -I^M, \quad (6)$$

$$\epsilon_{\text{L}}^{M-1} = -A^{M-1}, \quad (7)$$

and

$$\epsilon_{\text{H}}^M = \epsilon_{\text{L}}^{M-1}, \quad (8)$$

where the former two conditions have been expressed as Koopmans-type relationships. For each of these criteria, a tuning approach can be defined as a minimisation, on a system-by-system basis, of a norm measuring the extent to which that condition is satisfied. The norms based on the three tuning criteria above are denoted

$$H_{\text{LHS}} = |\epsilon_{\text{H}}^M + I^M|, \quad (9)$$

$$L_{\text{LHS}} = |\epsilon_{\text{L}}^{M-1} + A^{M-1}|, \quad (10)$$

$$\Omega_{\text{LHS}} = |\epsilon_{\text{L}}^{M-1} - \epsilon_{\text{H}}^M|, \quad (11)$$

respectively. We can similarly define the RHS analogues,

$$H_{\text{RHS}} = |\epsilon_{\text{H}}^{M+1} + I^{M+1}|, \quad (12)$$

$$L_{\text{RHS}} = |\epsilon_{\text{L}}^M + A^M|, \quad (13)$$

$$\Omega_{\text{RHS}} = |\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^{M+1}|. \quad (14)$$

In both cases, H and L refer to the Koopmans conditions appropriate to the given E vs. N segment, whilst Ω gives an indication of the degree of E vs. N curvature. It is worth noting that tuning to H or L constrains the orbital energies to be close to the *calculated* ionisation potential or electron affinity, but says nothing about how well they reproduce the exact quantities. Similarly, tuning to Ω constrains the two orbital energies/slopes to be equal to each other, but not necessarily equal to

the correct value. We shall investigate both of these issues in the present study.

By choosing one of the above minimisation criteria, one attempts to impose linearity on a single E vs. N segment. Baer and co-workers, however, noted^{32,38} that in order to obtain an accurate estimation of properties such as the fundamental gap, $I_0^M - A_0^M$, as a difference of Kohn–Sham orbital energies, *both* segments needed to be accurately described. To achieve this, they introduced two new tuning norms, which in our notation are written

$$J_{H,L} = H_{LHS} + L_{RHS}, \quad (15)$$

and

$$J_{H,H} = H_{LHS} + H_{RHS}. \quad (16)$$

By analogy, we can also define

$$J_{L,H} = L_{LHS} + H_{RHS}, \quad (17)$$

$$J_{L,L} = L_{LHS} + L_{RHS}. \quad (18)$$

$$(19)$$

A similar form for the tuning norm has also been proposed,^{35,38} differing from eq. 16 only in the cross-terms of its square, which we write as

$$\|J_{H,H}\|_2 = \sqrt{H_{LHS}^2 + H_{RHS}^2}, \quad (20)$$

and for which we can also consider the related combinations $\|J_{H,L}\|_2$, $\|J_{L,H}\|_2$, and $\|J_{L,L}\|_2$. Indeed, we observe that eqs. 16 and 20 are the first two terms of a series of p -norms,

$$\|J_{H,H}\|_p = \sqrt[p]{H_{LHS}^p + H_{RHS}^p}, \quad (21)$$

which leads us to consider the general p -norm

$$\|J_{x,y}\|_p = \sqrt[p]{x^p + y^p}, \quad (22)$$

where x and y refer to LHS and RHS conditions, respectively; we consider up to $p = 4$. Overall, this leads to a total of 22 tuning norms, listed in Table 1.

2.2 Assessment of tuning methods

We considered the same set of atoms as Baer and co-workers,^{32,38} comprising Li–F, Na–Cl and Ga–Br. The quantity M in the tuning norms is the number of electrons in the neutral atom. The ground state spin configuration was used for both neutral atoms and ions. Tuning was carried out using a range-separated^{40–47} form of B3LYP,^{48–52} denoted LC-B3LYP using the notation of Ref. 43, with the fraction of exact exchange varying from 0 to 100 % (i.e., $\alpha = 0$, $\beta = 1$) at a rate defined by the range-separation parameter μ . For each atom, calculations were performed using a series of μ values, and the value that minimised each norm, denoted μ^* , was determined to 2 d.p. For comparison, calculations were also carried out using a representative GGA (PBE⁵³), a conventional hybrid functional (B3LYP) and non-tuned range-separated hybrids (BNL^{47,54} and LC-B3LYP, each with range-separation parameter $\mu = 0.4a_0^{-1}$). Calculations were performed using the Gaussian 09⁵⁵ and CADPAC⁵⁶ programs, with the aug-cc-pVTZ basis set.

Inevitably, use of these tuning methods increases the computational effort, since calculations must be performed over a range of μ values. In the present study—where multiple criteria are investigated, each with a (potentially) different minimum—we used a coarse grid of μ values, refining as necessary close to the minima. For a single tuning norm the procedure could be carried out more efficiently, but would still require a number of separate calculations (typically greater than 20).

We assess the tuning methods in several ways. To test the Koopmans relationships, we consider the deviation of ϵ_{H}^M and ϵ_{L}^M from the exact $-I_0^M$ and $-A_0^M$, respectively;⁵⁷ it is also pertinent to

consider the deviation of the calculated I^M and A^M (determined from integer energy differences) from I_0^M and A_0^M . The quantities Ω_{LHS} and Ω_{RHS} provide a measure of linearity for the individual segments and so we consider the deviation of these quantities from zero (by construction, Ω will be near-zero when it is successfully employed as the tuning norm, but this will not necessarily be the case for a general tuning norm). Finally, we consider the deviation of $\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$ from the fundamental gap $I_0^M - A_0^M$. Mean absolute deviations are presented in Table 1, determined by calculating individual absolute deviations for each atom using its corresponding μ^* value and then averaging over the set of atoms. The corresponding standard deviations are presented in the Supporting Information. For clarity, the results are divided into those determined using conventional functionals and those determined using tuned functionals with single-segment (H , L , and Ω) and double-segment (J) tuning.

2.2.1 Conventional functionals

First, consider the results in Table 1 determined using the PBE functional. As expected, the values of ϵ_{H}^M and ϵ_{L}^M differ significantly from $-I_0^M$ and $-A_0^M$, by 3 to 4 eV. By contrast, the directly computed I^M and A^M are more than an order of magnitude more accurate. The large values of 6 to 8 eV for the linearity measures Ω_{LHS} and Ω_{RHS} quantify the significant E vs. N curvature. As noted in Section 1, the errors in ϵ_{H}^M and ϵ_{L}^M are of opposite sign and so the deviation is amplified when the difference is used to compute the gap. It follows that the gaps $\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$ deviate by more than 7 eV from $I_0^M - A_0^M$. In the case of a GGA such as PBE, the difference between the fundamental gap and the associated orbital energy gap is approximately equal to the exact exchange–correlation integer discontinuity;^{10,19} the relationship between curvature and integer discontinuity has recently been discussed by Stein et al.⁵⁸

B3LYP performs slightly better than PBE for each of the problematic quantities, reflecting the reduction in convexity caused by the addition of some exact exchange, whilst maintaining the low deviations in I^M and A^M . The non-tuned range-separated hybrids BNL and LC-B3LYP show a further marked improvement in the problematic quantities, although the deviations remain non-

Table 1: Mean absolute deviations (in eV) in various quantities, over all atoms, relative to exact reference values,⁵⁷ for conventional functionals and each tuning norm.

	ϵ_{H}^M	ϵ_{L}^M	I^M	A^M	Ω_{LHS}	Ω_{RHS}	$\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$
Conventional functionals							
PBE	4.10	3.04	0.15	0.19	8.22	5.71	7.13
B3LYP	3.16	2.35	0.17	0.17	6.57	4.37	5.50
BNL	0.76	0.33	0.15	0.32	1.46	0.50	0.85
LC-B3LYP	0.36	0.37	0.26	0.24	1.29	0.31	0.62
Single-segment tuning							
H_{LHS}	0.26	0.33	0.26	0.17	0.16	0.79	0.53
L_{LHS}	0.34	0.40	0.25	0.16	0.17	0.91	0.68
Ω_{LHS}	0.30	0.37	0.26	0.17	0.09	0.85	0.61
H_{RHS}	0.63	0.42	0.26	0.23	1.82	0.21	0.92
L_{RHS}	0.30	0.22	0.26	0.21	1.16	0.21	0.39
Ω_{RHS}	0.46	0.32	0.25	0.23	1.45	0.04	0.65
Double-segment tuning							
$\ J_{\text{H,H}}\ _1$	0.24	0.32	0.26	0.17	0.19	0.78	0.51
$\ J_{\text{H,H}}\ _2$	0.09	0.21	0.26	0.19	0.64	0.49	0.19
$\ J_{\text{H,H}}\ _3$	0.10	0.20	0.26	0.19	0.71	0.45	0.17
$\ J_{\text{H,H}}\ _4$	0.11	0.20	0.27	0.19	0.73	0.44	0.16
$\ J_{\text{H,L}}\ _1$	0.25	0.32	0.26	0.17	0.18	0.78	0.52
$\ J_{\text{H,L}}\ _2$	0.17	0.26	0.26	0.18	0.37	0.65	0.34
$\ J_{\text{H,L}}\ _3$	0.16	0.24	0.26	0.18	0.42	0.62	0.31
$\ J_{\text{H,L}}\ _4$	0.14	0.23	0.27	0.18	0.45	0.61	0.28
$\ J_{\text{L,H}}\ _1$	0.33	0.39	0.25	0.16	0.16	0.89	0.67
$\ J_{\text{L,H}}\ _2$	0.11	0.22	0.26	0.19	0.55	0.55	0.22
$\ J_{\text{L,H}}\ _3$	0.11	0.20	0.26	0.19	0.61	0.51	0.19
$\ J_{\text{L,H}}\ _4$	0.11	0.20	0.26	0.19	0.64	0.50	0.17
$\ J_{\text{L,L}}\ _1$	0.34	0.40	0.25	0.16	0.17	0.90	0.67
$\ J_{\text{L,L}}\ _2$	0.22	0.29	0.26	0.18	0.26	0.72	0.44
$\ J_{\text{L,L}}\ _3$	0.19	0.27	0.26	0.18	0.33	0.68	0.38
$\ J_{\text{L,L}}\ _4$	0.18	0.26	0.26	0.18	0.35	0.67	0.37
Reference value	$-I_0^M$	$-A_0^M$	I_0^M	A_0^M	0	0	$I_0^M - A_0^M$

negligible. With these two functionals, the I^M and A^M values degrade marginally.

2.2.2 Single-segment tuning

Next consider the functionals tuned to criteria on a single E vs. N segment, using H , L or Ω as tuning norms. The deviations in ϵ_{H}^M and ϵ_{L}^M are generally close to those from the non-tuned range-separated functionals. The lowest deviations in ϵ_{H}^M and ϵ_{L}^M are, not surprisingly, obtained by tuning to H_{LHS} and L_{RHS} , respectively, since these explicitly optimise the Koopmans conditions that are being assessed. Importantly, these deviations are still non-zero, challenging the assumption that simply tuning to the calculated I^M or A^M is sufficient—one must also consider the quality of the calculated I^M or A^M itself, i.e., how accurate the relative energies of the integer-electron systems are. In fact, I^M is in virtually constant deviation by 0.25 eV whichever tuning is used, and—somewhat counterintuitively— A^M is slightly better when tuning the LHS rather than the RHS.

Tuning to H_{LHS} and L_{LHS} gives comparatively low values of Ω_{LHS} , up to five times smaller than Ω_{RHS} , indicating that the tuning is relatively successful at linearising the LHS segment, but at the expense of the RHS. Tuning to H_{RHS} and L_{RHS} yields analogous behaviour, with Ω_{RHS} values up to nine times smaller than Ω_{LHS} . When used as tuning norms, Ω_{LHS} and Ω_{RHS} give, by construction, near-zero values for their respective linearity measures (although large values are seen on the non-optimised side). However, all that a small value of Ω indicates is that the slopes at the two integers are essentially the same—not necessarily that they are equal to the correct value. In fact, the discrepancy between the deviations in ϵ_{H}^M and ϵ_{L}^M and those in the calculated I^M and A^M confirms that the slopes are not correct; see Section 2.3.

For both LHS and RHS tuning norms, the deviations in the gap $\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$ are of a similar magnitude to the non-tuned range-separated functionals. The lowest deviations are obtained when tuning to H_{LHS} and L_{RHS} , reflecting the fact that these two conditions individually yield the most accurate ϵ_{H}^M and ϵ_{L}^M , respectively, which are the two components of the gap.

2.2.3 Double-segment tuning

Next consider the functionals tuned to criteria on both E vs. N segments, using the various J in Eq. (22) as tuning norms. In all cases, the deviations in ϵ_{H}^M and ϵ_{L}^M reduce notably from $p = 1$ to $p = 2$, with little subsequent change for $p > 2$; the best results are obtained using the $\|J_{\text{H,H}}\|_p$ series, with deviations as small as 0.1 to 0.2 eV. However, similar trends are not seen in the computed I^M and A^M —the deviation in I^M is an almost constant 0.26 eV on average, for all of the tuning norms tested. This has obvious implications for the tuning methods, which rely on attempting to constrain the frontier orbital energies to these incorrect I^M values.

As p increases beyond unity, Ω_{LHS} increases, which is somewhat counterintuitive given that ϵ_{H}^M improves. A more intuitive trend is observed for Ω_{RHS} , which reduces as ϵ_{L}^M improves. Despite yielding the most accurate ϵ_{H}^M and ϵ_{L}^M , the values of Ω_{LHS} and Ω_{RHS} are both significant for the $\|J_{\text{H,H}}\|_p$ series with $p > 1$, indicating that substantial non-linearity remains. Insight into these observations is provided in Section 2.3.

The accuracy of the gaps $\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$ again reflect the accuracy of the orbital energies: the best results are obtained for the $\|J_{\text{H,H}}\|_p$ series, with deviations reducing to less than 0.2 eV.

2.3 A representative system: the carbon atom

Further insight into the behaviour of the tuned functionals is obtained by focusing on a single atom, thereby allowing the quantities in Table 1 to be explicitly related to an E vs. N curve. We consider the carbon atom with $M = 6$ and $5 \leq N \leq 7$. Table 2 presents μ^* values determined for the carbon atom, using selected tuning norms, along with the deviations (calculated minus reference) in ϵ_{H}^M , ϵ_{L}^M , I^M , A^M , Ω_{LHS} , Ω_{RHS} , and $\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$, computed using μ^* . The dependence of these quantities on the choice of tuning norm largely follows the behaviour of the average quantities in Table 1.

For each tuning norm in Table 2, an E vs. N curve was produced by fixing μ at the corresponding value of μ^* and smoothly varying the number of α electrons from 3 to 5, with the number of β electrons fixed at 2. To most effectively illustrate the non-linearity of the E vs. N curve, we plot the deviation of the calculated energy from a linear interpolation between the calculated energies

at integer N ; we will refer to these curves as E vs. N deviation curves. By construction, the interpolated and calculated energies agree at integer. For non-integer values, a horizontal line along zero indicates a linear E vs. N curve, whilst a positive/negative deviation indicates a concave/convex curve. Results are presented in Figures 2-4.

A pair of straight lines is superimposed onto each E vs. N deviation curve, one on the electron-deficient side and one on the electron-abundant side of $N = 6$. These lines are the differences between the *exact* piecewise curve and the linear interpolation between calculated integers, aligned at $N = 6$. The slopes of the lines indicate the limiting slopes that an E vs. N deviation curve would have to exhibit at $N = 6$, in order to yield $\epsilon_{\text{H}}^M = -I_0^M$ and $\epsilon_{\text{L}}^M = -A_0^M$, respectively. We term them ‘exact slopes’ in the context of an E vs. N deviation plot. These exact slopes provide a useful guide to the quality of a functional: the difference between the slope of the E vs. N deviation curve and the exact slope quantifies the deviations in ϵ_{H}^M and ϵ_{L}^M from $-I_0^M$ and $-A_0^M$, whilst the deviation of the exact slope from a horizontal line quantifies the deviations in I^M and A^M , again from $-I_0^M$ and $-A_0^M$. We note that if the exact slope is not horizontal then satisfaction of the exact Koopmans conditions ($\epsilon_{\text{H}}^M = -I_0^M$; $\epsilon_{\text{L}}^M = -A_0^M$) will *require* non-linearity in the E vs. N curve.

Figure 2 presents the E vs. N deviation curves for the three μ^* values determined by tuning to the RHS. The three μ^* values are rather different from one another and so the E vs. N behaviour of each is also quite different. In moving from H_{RHS} to Ω_{RHS} to L_{RHS} , the slopes on either side of $N = 6$ move closer to the exact slopes, and so the deviations in ϵ_{H}^M and ϵ_{L}^M in Table 2 reduce. The exact slopes are notably offset from horizontal, reflecting the deviations in I^M and A^M in Table 2. By construction, tuning to H_{RHS} and L_{RHS} yields—for the RHS segment—near-zero slopes at $N = 7$ and $N = 6$, respectively. However, in both cases the unconstrained end of the RHS segment exhibits a much larger slope, leading to the small but non-negligible Ω_{RHS} values in Table 2; by contrast the LHS segment is highly convex, with large Ω_{LHS} values. By construction, tuning to Ω_{RHS} yields essentially identical slopes at $N = 6$ and $N = 7$ for the RHS segment and hence near-zero values for Ω_{RHS} , but the slopes themselves are not zero, resulting in a curve with a point of inflection. The LHS segment is again highly convex, with a correspondingly large Ω_{LHS} value.

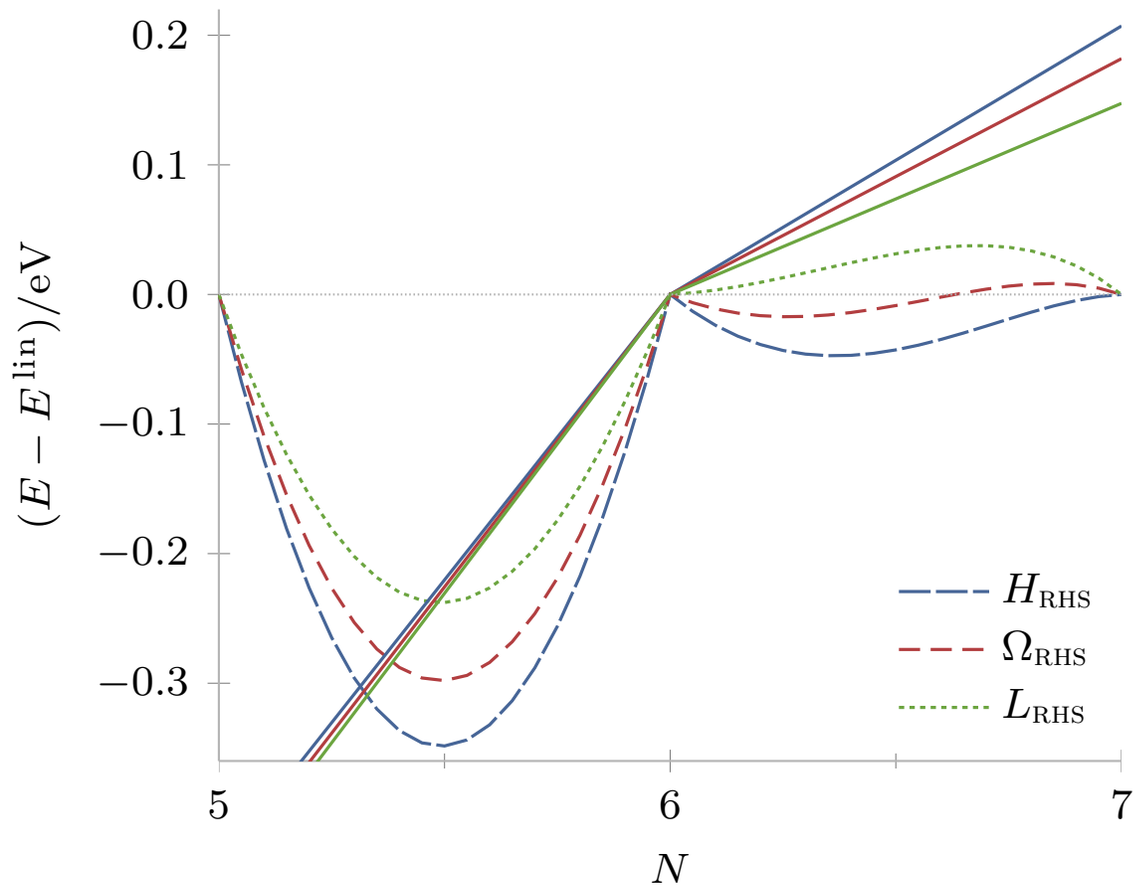


Figure 2: E vs. N deviation curves (dashed/dotted curves) and exact slopes (solid straight lines) for the carbon atom using RHS tuning norms. See text for definitions of these quantities.

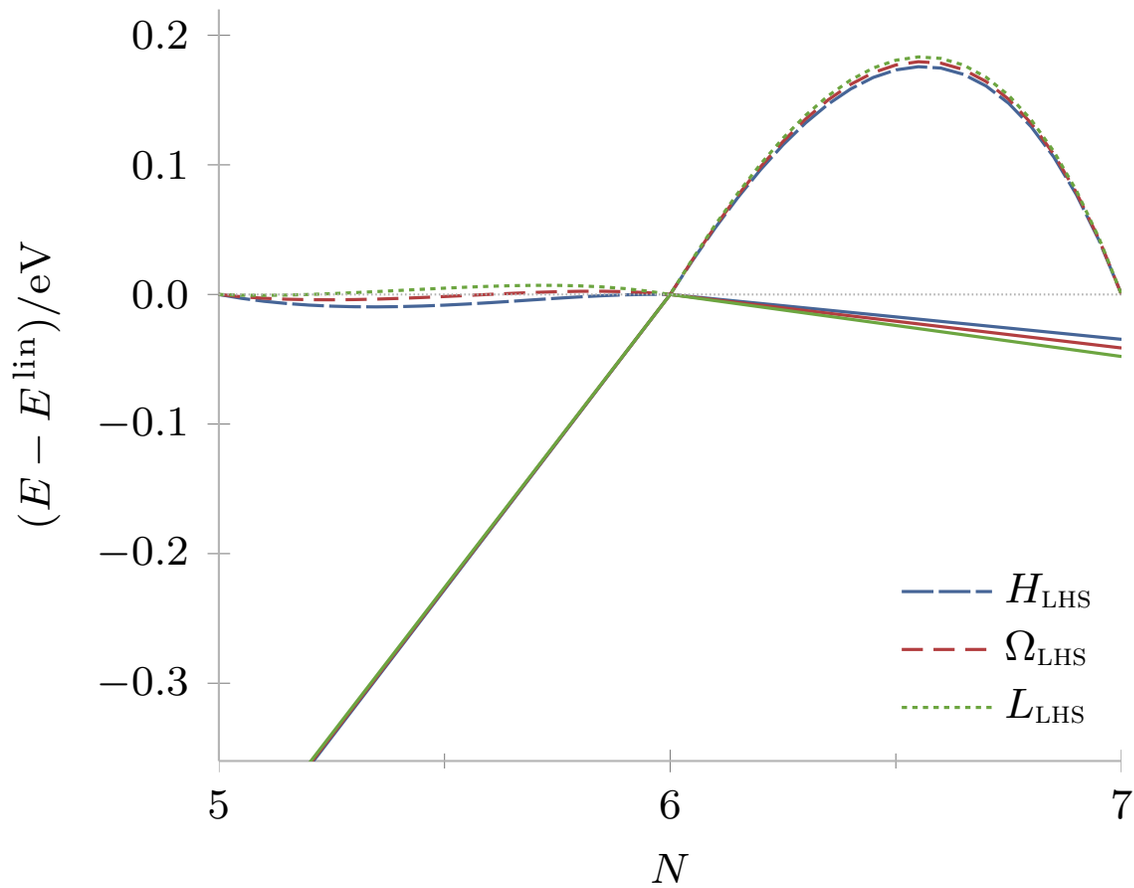


Figure 3: E vs. N deviation curves (dashed/dotted curves) and exact slopes (solid straight lines) for the carbon atom using RHS tuning norms. See text for definitions of these quantities.

Table 2: Deviations (in eV) in various quantities, for the carbon atom, relative to exact reference values,⁵⁹ for each tuning norm. μ^* is the optimal range-separation parameter (in a_0^{-1}).

	μ^*	ϵ_{H}^M	ϵ_{L}^M	I^M	A^M	Ω_{LHS}	Ω_{RHS}	$\epsilon_{\text{L}}^M - \epsilon_{\text{H}}^M$
Single-segment tuning								
H_{LHS}	0.66	-0.47	0.57	0.46	-0.03	-0.05	1.44	1.03
L_{LHS}	0.68	-0.52	0.61	0.45	-0.05	0.05	1.50	1.12
Ω_{LHS}	0.67	-0.49	0.59	0.45	-0.04	0.00	1.47	1.08
H_{RHS}	0.36	0.89	-0.50	0.44	0.21	-2.75	-0.31	-1.38
L_{RHS}	0.43	0.44	-0.14	0.46	0.15	-1.87	0.29	-0.57
Ω_{RHS}	0.39	0.68	-0.33	0.45	0.18	-2.35	-0.03	-1.01
Double-segment tuning								
$\ J_{\text{H,H}}\ _1$	0.65	-0.44	0.55	0.46	-0.03	-0.11	1.41	0.99
$\ J_{\text{H,H}}\ _2$	0.53	-0.04	0.24	0.47	0.06	-0.91	0.91	0.29
$\ J_{\text{H,H}}\ _3$	0.52	0.00	0.21	0.47	0.07	-1.00	0.86	0.21
$\ J_{\text{H,H}}\ _4$	0.51	0.04	0.18	0.47	0.08	-1.08	0.81	0.14
Reference value		$-I_0^M$	$-A_0^M$	I_0^M	A_0^M	0	0	$I_0^M - A_0^M$

Figure 3 shows the analogous curves when tuning to the LHS. The three μ^* values are now very close to one another and so the differences in the E vs. N behaviour are much less pronounced. The near-linearity of the LHS is much more pronounced than the RHS was in Figure 2 and so, at first sight, one might expect an accurate ϵ_{H}^M . However, the plot simply illustrates that $\epsilon_{\text{H}}^M \approx -I^M$; by contrast the deviation of the exact slope from horizontal indicates that $I^M \neq I_0^M$ and so the discrepancy between ϵ_{H}^M and $-I_0^M$ is actually significant. The deviation in ϵ_{L}^M is of a similar magnitude—in this case, the deviation arises largely due to the curvature, rather than the error in A^M , which is now much smaller.

Finally, consider the curves obtained by tuning to both segments. Each set of p -norms shows a similar trend, so we choose the most successful method, $\|J_{\text{H,H}}\|_p$, to illustrate the behaviour. Figure 4 presents the E vs. N deviation curves for the μ^* values obtained for $1 \leq p \leq 4$. When $p = 1$, $\mu^* = 0.65 a_0^{-1}$, which is essentially the same as the value obtained when tuning to the LHS segment, and so the curve is close to those in Figure 3. Increasing p to 2 yields $\mu^* = 0.53 a_0^{-1}$, roughly midway between the LHS-only and RHS-only optimised values. This results in some

reduction in concavity on the RHS (Ω_{RHS} in Table 2 decreases) but increased convexity on the LHS (Ω_{LHS} increases in magnitude), so that neither side shows near-linear behaviour. As p is increased to 3 and 4, μ^* decreases marginally again, with a further small shift in the E vs. N deviation curves. Despite the lack of linearity on either side, using $p > 1$ yields slopes that are closest to the exact slopes, and hence the values of ε_{H}^M and ε_{L}^M are optimal. The good performance of the $p > 1$ functionals therefore arises from a convenient error cancellation between lack of linearity and errors in I^M and A^M .

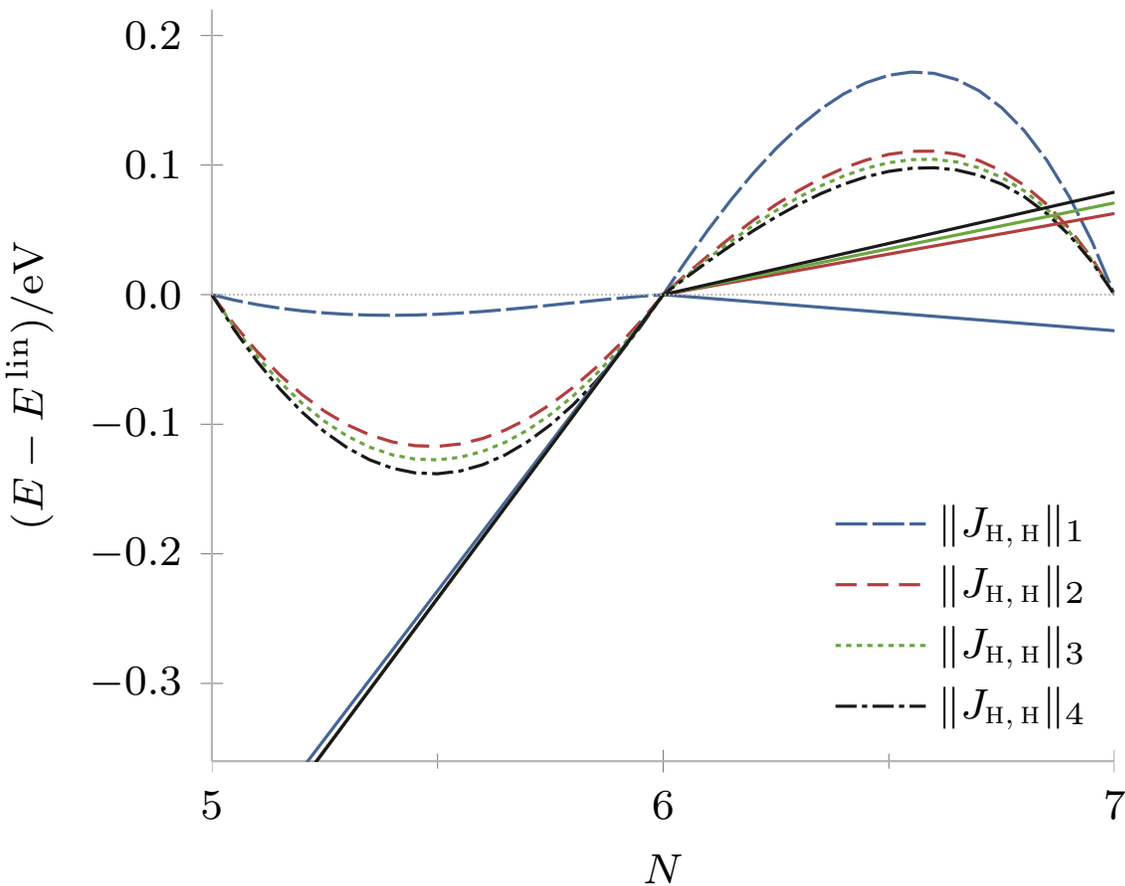


Figure 4: E vs. N deviation curves (dashed/dotted curves) and exact slopes (solid straight lines) for the carbon atom using RHS tuning norms. See text for definitions of these quantities.

3 Conclusions

We have assessed a range of tuning methods for enforcing approximate energy linearity, i.e., reducing delocalisation error, through a system-by-system optimisation of a range-separated hybrid functional. For a series of atoms, the accuracy of frontier orbital energies, ionisation potentials, electron affinities, and orbital energy gaps has been quantified and particular attention has been paid to the extent to which approximate linearity is actually achieved in the resulting E vs. N curve.

The tuning approaches can yield significantly improved orbital energies and orbital energy gaps, compared to those from conventional functionals. For M -electron systems, optimal results were obtained using a tuning norm based on the HOMO energy of the M and $M + 1$ electron systems, with deviations of just 0.1 to 0.2 eV in these quantities, compared to exact values. However, detailed examination for the carbon atom illustrates a subtle cancellation between errors arising from non-linearity and errors in the computed ionisation potentials and electron affinities used in the tuning.

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5 Supporting Information

Standard deviations corresponding to the data in Table 1 and E vs. N curves corresponding to the data in Figures 2-4.

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