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## Extensive Reference Set and Refined Computational CP000431K Protocol for Calculations of <sup>57</sup>Fe Mössbauer Parameters

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**ORCIDs**:

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ABSTRACT

View Article Online
DOI: 10.1039/D4CP00431K

Mössbauer spectroscopy is a powerful technique for probing the local electronic structure of iron compounds, because it reports in an element-selective manner on both the oxidation state and coordination environment of the Fe ion. Computational prediction of the two main Mössbauer parameters, isomer shift ( $\delta$ ) and quadrupole splitting ( $\Delta E_0$ ), have long been targeted by quantum chemical studies, and useful protocols based on density functional theory have been proposed. Here we present an extensive curated reference set of Fe compounds that is considerably larger and more diverse than literature precedents. We make a distinction between low-temperature and high-temperature experimental subgroups. This set is employed for optimizing a refined computational protocol utilizing the scalar version of the exact 2component (X2C) Hamiltonian with the finite nucleus approximation. Attention is devoted to having an accurate and flexible all-electron basis set for Fe. We assess the performance of several DFT methods that cover all representative families and rungs of functionals and find that hybrid functionals with ca. 25-30% exact exchange offer the best accuracy for isomer shifts. The work establishes a refined general protocol of wide applicability that achieves good performance for the prediction of isomer shifts in a wider variety of systems than before, but the limitations of DFT for quadrupole splittings are also highlighted. Finally, comparison of calculated values with high-temperature experimental results shows that the use of an empirical correction factor is required to account for the second-order Doppler shift and to achieve the same quality of correlation as with the low-temperature data.

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1. Introduction

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After the discovery of the Mössbauer effect in 1958,<sup>1-3</sup> Mössbauer spectroscopy has emerged as a powerful analytic tool in solid state physics, chemistry, metallurgy, biological- and geosciences as well as in industrial and materials science applications. This spectroscopic technique is based on the phenomenon of recoilless resonant absorption of gamma-ray photons by an atomic nucleus and it can successfully probe minute changes in the nuclear energy levels originating from hyperfine interactions of the active center with surrounding electrons. Over the years, the Mössbauer effect has been observed for nearly 90  $\gamma$ -ray transitions in 72 isotopes of 42 different elements.<sup>4</sup> However, <sup>57</sup>Fe Mössbauer spectroscopy is the most common due to the importance of iron in homogeneous and heterogeneous catalysis as well as the abundance of different spin and oxidation states of this element in biologically relevant systems.<sup>5-9</sup>

The two most important parameters encoded in a Mössbauer spectrum are the isomer shift ( $\delta$ ) and the quadrupole splitting ( $\Delta E_{\rm Q}$ ). The isomer shift of a metal center is directly related to the electron density at the nucleus, while  $\Delta E_{\rm Q}$  is proportional to the electric field gradient (EFG), which originates from a nonsymmetrical distribution of electrons in the valence shell as well as charges on the neighboring ligands.

The isomer shift of  $^{57}$ Fe measures the shift in the energy of the  $\gamma$ -ray absorption relative to a standard, usually iron foil. The isomer shift is sensitive to the electron density at the nucleus, and indirectly probes changes in iron-ligand bond lengths, covalency and nature of its bonds, and shielding due to the 3d orbital occupation pattern. As a result, it can successfully probe oxidation and spin states, and the coordination environment of Fe. This shift between an absorber (A) and a source (S) comes from the difference in the electrostatic interactions between electronic and nuclear charge distributions, which originates from the difference in their electron densities as well as the change in the nuclear radius upon gamma transition. Considering the nucleus to be a uniformly charged sphere, the mathematical expression for the Mössbauer isomer shift is

$$\delta = \alpha \left( \overline{\rho}_{e}^{A} - \overline{\rho}_{e}^{S} \right) \tag{1}$$

where  $\overline{\rho}_e$  denotes the "effective" electron density, i.e., the weighted average of the electronicle Online density within the finite volume the nucleus;  $\alpha$  is the isomer shift "calibration constant" and can be expressed with the following form

$$\alpha = \frac{4\pi cZ}{5E_{\gamma}} R_0^2 \left(\frac{\Delta R_0}{R_0}\right) \tag{2}$$

where c,  $E_{\gamma}$ ,  $R_0$ , and Z are the speed of light, energy of the gamma quantum, radius of the nucleus, and the nuclear charge, respectively. The  $\Delta R_0/R_0$  ratio in Eq. (2) describes the relative change of the nuclear radius upon excitation. For a specific Mössbauer isotope, with the exception of  $\overline{\rho}_e^A$  all terms in Eq. (1) are constant. In the standard computational approach, the "effective" electron density is usually approximated by the contact density,  $\rho(0)$ . Although for lighter elements like <sup>57</sup>Fe this approximation produces negligible error, for heavier nuclei this error is nontrivial. Considering this approximation, Eq. (1) can be written in a simple form as

$$\delta = a[\rho(0) - C] + b \tag{3}$$

where a and b are the "fitting" or "correlation" constants, which can be determined from a linear fit of the computed  $\rho(0)$  vs. experimental isomer shifts ( $\delta_{\rm exp}$ ) for a set of iron complexes. These fitting constants contain not only nuclear information, but also cover for the deficiencies of the quantum-chemical treatment. Hence, the fitting constant "a" is different from the calibration constant  $\alpha$  defined in Eq. (1). The constant, C is introduced only for a convenient scaling of the very large values of contact densities. The success of such ansatz to predict good correlation between theory and experiment relies to some extent on error cancellation<sup>11</sup> and suffers from the fact that each new combination of quantum chemical protocol would require new correlation constants. We should note that there is a physically "correct" slope of the correlation line, with  $\alpha = -0.31 \pm 0.04$ .

Over the years, linear regression analysis has been extensively applied for the calculation of isomer shifts using semi-empirical, Hartree–Fock (HF), density functional theory (DFT), and wave function based ab initio methods like the domain based local pair-natural orbital coupled-cluster theory (DLPNO-CCSD).<sup>11, 13-34</sup> All these studies have demonstrated good correlation between theory and experiment. Despite its simplicity, this approach is fairly reliable and

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efficient and is known to predict isomer shifts with an accuracy of up to  $\sim$ 0.1 mm s<sup>-1.27</sup> Accirrate colling computation of contact density is challenging and depends on factors like the choice of the quantum chemical method, the basis set, and the proper treatment of relativistic effects. An ideal basis set which can adequately describe the contact density of Fe has to be sufficiently large in the region where a cusp in the electron density will occur. The known nonrelativistic HF limit for  $\rho(0)$  is  $\sim$  11,903.987 a.u.<sup>-3</sup>, whereas with a good basis set one can only obtain up to 11,820 a.u.<sup>-3</sup>.16, 19 Although in terms of absolute contact density this error is negligible, compared to the variation of the electron density over the chemical range ( $\sim$ 10 a.u.<sup>-3</sup>) it is nontrivial.

The relativistic effects on the electron density at the nucleus are large for iron. It is known to increase  $\rho(0)$  by a factor of 1.3 compared to the nonrelativistic electron density of  $^{57}$ Fe. $^{17, 19}$  On the other hand, Saue and co-workers have shown that for the calculation of Mössbauer isomer shifts spin-orbit coupling can be safely ignored while using the eXact 2-Component (X2C) Hamiltonian $^{35-38}$  and a finite nucleus model, $^{39}$  therefore the consideration of scalar relativistic effects is sufficient. $^{40}$  As already demonstrated, $^{11}$  the contributions from the core 1s and 2s orbitals remain nearly constant to the absolute contact density at the iron nucleus. The major contribution to the variation of the contact density due to its electronic configuration and ligand environment arise from the valence and subvalence regions. Hence, these inner valence and outer valence orbitals of iron matter the most for obtaining a correct calibration. $^{11, 19}$ 

The other important parameter in a Mössbauer spectrum is the quadrupole splitting ( $\Delta E_Q$ ), which arises from the interaction of the nuclear quadrupole moment of the excited state with the EFG at the nucleus. Quadrupole splittings can also be used as a sensitive probe for the coordination environment of iron centers. The quadrupole splitting is obtained from the EFGs using the expression

$$\Delta E_{Q} = \frac{1}{2} e Q V_{zz} \sqrt{1 + \frac{1}{3} \left\{ \frac{|V_{yy} - V_{xx}|}{V_{zz}} \right\}^{2}}$$
 (4)

where e is the elementary charge and Q denotes the nuclear quadrupole moment (in barn, b, where 1 barn =  $10^{-28}$  m<sup>2</sup>), and  $V_{xx}$ ,  $V_{yy}$ , and  $V_{zz}$  indicate the components of the EFG tensor in the

principal axis system  $\{x, y, z\}$ . Although DFT has become the workhorse for the prediction of fice online isomer shifts with reasonable success, the predictions of quadrupole splittings are often associated with larger errors. This is presumably because of the inability of such calculations to represent the finer details of the asymmetry in electron density.

Another important factor is the absence of an accurate value of the  $^{57}$ Fe nuclear quadrupole moment (NQM). As the NQM is impossible to determine experimentally, the only possible way to obtain those values is via linear regression analysis using the experimental  $\Delta E_{\rm Q}$  values and the theoretical EFGs. Therefore, a wide range of values between 0.1 to 0.3 b can be found in the literature. In the present study, we have used Q = 0.160 b for the calculation of quadrupole splittings.<sup>17</sup>

Over the years, several benchmark studies on  $^{57}$ Fe Mössbauer parameters have been performed by employing scalar relativistic Hamiltonians like ZORA (zero-order regular approximation),  $^{41}$  DKH2 (second-order Douglas–Kroll–Hess),  $^{42-48}$  and X2C (exact two-component).  $^{18, 19, 21, 22, 40, 49}$  Although changing the Hamiltonian had little effect on the overall correlation of experimental  $\delta$  and calculated contact densities of iron,  $^{18, 22}$  a detailed evaluation of a variety of density functional methods and basis sets with scalar relativistic X2C Hamiltonian against a large and diverse dataset is still absent. Another potential constraint of previous studies is related to the set of iron complexes considered for benchmarking quantum-chemistry methods. These datasets are either composed of limited number of compounds, a mixture of molecular and solid-state systems, a specific type of ligand in focus, or only a limited number of spin and oxidation states of iron. The main objectives of the present study are:

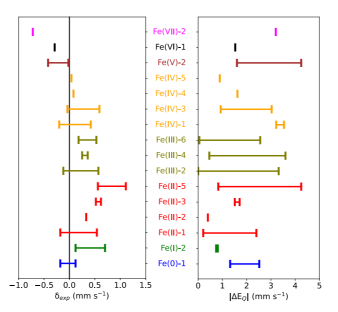
- (a) To construct a complete database of Fe complexes with well-established Mössbauer parameters, which is curated so that it includes most of the known spin and oxidation state of iron and is representative of the wide range of chemical types encountered in iron coordination chemistry;
- (b) To conduct a thorough benchmark study of basis sets and DFT functionals against our new database in order to develop a refined computational protocol based on the popular X2C Hamiltonian.

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Starting from 155 iron-containing systems with one Fe center, 116 unique entries were sorted out depending on the nature of ligands, spin and oxidation states of the Fe center, and distribution of the experimental isomer shift ( $\delta_{\rm exp}$ ) and quadrupole splitting ( $\Delta E_{\rm Q}$ ) values. Among these, 101 are molecular iron complexes, where 80 experimental isomer shifts were recorded at low temperature (4.2–80 K) and the remaining 21 were recorded at –or close to–room temperature. In the following, we shall focus on the subset of 80 molecular iron-complexes for which low-T  $\delta_{\rm exp}$  values are known. We call this dataset MPMIC80 ( $\underline{\bf M}$  össbauer  $\underline{\bf P}$  arameters of 80  $\underline{\bf M}$  olecular Iron  $\underline{\bf C}$  omplexes). Among these complexes,  $\delta_{\rm exp}$  ranges from –0.72 to 1.11 mm s<sup>-1</sup>, and the absolute quadrupole splitting ( $|\Delta E_{\rm Q}|$ ) values range from 0.0 to 4.25 mm s<sup>-1</sup> (see Figure 1). The complexes in our dataset cover the known oxidation and spin states of iron. For detailed information on spin, oxidation states,  $\delta_{\rm exp}$ ,  $|\Delta E_{\rm Q}|$ , and corresponding references see Table S1 in the ESI.



**Figure 1.** Distribution of experimental isomer shifts  $\delta_{\text{exp}}$  (mm s<sup>-1</sup>) and quadrupole splitting values  $|\Delta E_{\text{Q}}|$  (mm s<sup>-1</sup>) for different oxidation states of Fe and total spin multiplicities (2*S*+1) of the complexes present in the MPMIC80 data set.

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## 3. Methodology

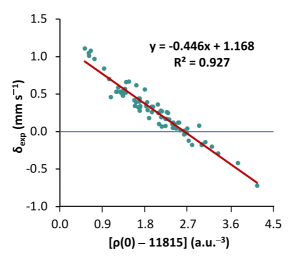
All the calculations were carried out using a development version of ORCA.<sup>50</sup> All molecular iron complexes were optimized at the gas-phase using the scalar relativistic X2C Hamiltonian,<sup>51</sup> the TPSSh<sup>52, 53</sup> functional, the D3BJ dispersion corrections, <sup>54, 55</sup> and the x2c-TZVPall basis sets.<sup>56</sup> During optimization, the DefGrid2 integration grid, TightSCF convergence criteria, RIJCOSX approximation,<sup>57</sup> and picture change effects<sup>58-61</sup> were included. Using two selected complexes, we validated that DefGrid2 is already fully converged for optimizations with the above-mentioned methodology (see Table S3). For the scalar relativistic calculations of Mössbauer parameters, we used the x2c-TZVPPall<sup>56</sup> basis set for the ligand atoms, we employed first-order picture change effects and the finer DefGrid3 integration grids throughout. In the present study the performance of 10 different basis sets for iron is evaluated: CP(PPP), <sup>11</sup> x2c-TZVPPall,<sup>56</sup> x2c-TZVPPall-s,<sup>62</sup> aug-cc-pVTZ-J (or aVTZ-J),<sup>63</sup> DKH-def2-TZVPP (exponents from def2-TZVPP<sup>64</sup> were recontracted for scalar-relativistic DKH2<sup>43-47, 65</sup> Hamiltonian), ANO-RCC-VTZP,66 s-decontracted x2c-TZVPPall, s-decontracted aug-ccpVTZ-J, s-decontracted aug-cc-pVTZ-J(-dfg), and aug-cc-pVTZ-Jmod.<sup>67</sup> The last basis set was proposed by Gómez-Piñeiro et al. for the calculations of Cu(II) core properties, where the augcc-pVTZ-J was modified by decontracting the s functions and removing the three innermost primitives.<sup>67</sup> On the other hand, the outermost d-, f- and g-primitives are removed from sdecontracted aug-cc-pVTZ-J to obtain the s-decontracted aug-cc-pVTZ-J(-dfg) basis set. For the ligand atoms, x2c-TZVPPall<sup>56</sup> was used throughout. 15 different density functionals from all five rungs of Jacob's ladder<sup>68</sup> were calibrated: SVWN5,<sup>69, 70</sup> BP86,<sup>71, 72</sup> PBE,<sup>73</sup> BLYP,<sup>72, 74</sup> TPSS,<sup>53</sup> PBE0,<sup>75</sup> B1LYP,<sup>76</sup> B3LYP,<sup>74, 77, 78</sup> TPSSh,<sup>52, 53</sup> TPSS0,<sup>79</sup> M06,<sup>80</sup> LC-BLYP,<sup>81</sup> CAM-B3LYP,<sup>82</sup> ωB97X,<sup>83</sup> and B2PLYP.<sup>84</sup> For the PT2 part of the double hybrid functionals, we correlated all core electrons and employed both relaxed and unrelaxed densities. The ORCA sample input files for the calculation of Mössbauer properties and the modified basis sets are provided in the ESI. In a previous study, the accuracy of selected DFT functionals was evaluated on a small set of 20 iron-containing compounds. 18 The structures were optimized

there using the TPSS functional and def2-TZVP85 basis set. Following the same protocol for

## 4. Results and discussion

## 4.1. Isomer Shifts (ISs) on the MPMIC80 Set

First, to orient ourselves, linear regression analysis was performed for the calculated nonrelativistic electron densities using the originally proposed protocol<sup>18</sup> against the experimental isomer shifts of MPMIC80. The coefficient of determination ( $R^2$ ), a and b values obtained from the linear fit were 0.927, -0.446, and 1.168, respectively (see Figure 2).



**Figure 2.** Linear fitting of the nonrelativistic contact densities relative to the experimental isomer shifts of the 80 complexes in MPMIC80. The contact densities are obtained with the TPSS0 functional and the CP(PPP) basis set for Fe.

These results are worse than literature expectations,  $^{18, 22}$  owing to the considerably expanded reference set of compounds in the present study. The mean absolute error and standard deviation of calculated isomer shift ( $\delta_{cal}$ ) with respect to  $\delta_{exp}$  are 0.07 and 0.09 mm s<sup>-1</sup>, respectively. Closer inspection reveals that the calculated isomer shifts of all nine Fe(II) complexes with quintet spin multiplicity (2S+1=5) deviate significantly from the experimental data. This stresses the

necessity of reconsidering the linear correlation parameters obtained from more restricted the Online reference sets.

## 4.1.1 Basis set definition

Aiming to identify suitable basis sets to obtain a good correlation between  $\delta_{\rm exp}$  and  $\rho(0)$ , slope of the linear fit close to the experimentally determined "a" value, <sup>12</sup> and calculated contact density close to the four-component fully relativistic HF electron density of Fe(II)<sup>86</sup> we evaluated the performance of ten different basis sets for iron using the B3LYP functional, the Gaussian finite nucleus model of Visscher and Dyall, <sup>87</sup> and first-order picture change effects. The values of different fitting parameters and coefficient of determination ( $R^2$ ) obtained from the linear regression analysis, mean absolute deviation, standard deviation, and maximum deviation of  $\delta_{\rm cal}$  from  $\delta_{\rm exp}$  are listed in Table 1. For the results with the point nucleus model and picture change effect, see Table S4 in the ESI.

**Table 1**: Calibration constants and statistical parameters obtained from the linear fitting of Eq. (3) using ten different basis sets for iron and the B3LYP functional.<sup>[a, b]</sup>

Basis set for Fe	а	b	C	$\mathbb{R}^2$	MAD <sup>[c]</sup> (mm s <sup>-1</sup> )	Max. Dev. <sup>[d]</sup> (mm s <sup>-1</sup> )	St. Dev. <sup>[e]</sup> (mm s <sup>-1</sup> )
CP(PPP)	-0.30	0.9704	14362	0.956	0.054	0.191	0.070
x2c-TZVPPall	-0.22	1.0294	13651	0.820	0.108	0.346	0.142
x2c-TZVPPall-s	-0.22	0.8917	13652	0.806	0.113	0.356	0.147
aug-cc-pVTZ-J	-0.41	1.3456	10511	0.956	0.054	0.181	0.070
ANO-RCC-VTZP	-0.18	0.8903	15631	0.414	0.184	0.752	0.255
DKH-def2-TZVPP	-0.30	1.0315	14076	0.956	0.055	0.190	0.070
s-decontracted x2c-TZVPPall	-0.31	1.1020	13688	0.955	0.054	0.197	0.071
s-decontracted aug-cc-pVTZ-J	-0.29	1.1073	14930	0.958	0.053	0.180	0.069
s-decontracted aug-cc-pVTZ-J (-dfg)	-0.29	1.1051	14930	0.958	0.053	0.181	0.069
aug-cc-pVTZ-Jmod <sup>67</sup>	-0.29	1.1015	14804	0.958	0.053	0.180	0.069
Exp. <sup>12</sup>	-0.31±0.04						

<sup>[</sup>a] With first-order picture change effect and finite nucleus model. For the results obtained with the point nucleus model and picture change effect, see Table S4 in the ESI.

 $<sup>^{[</sup>b]}a$  and b are the fitting coefficients obtained from the linear fit of Eq. (3). C is a constant, which is very close to the calculated  $\rho(0)$  value. The units of a, b, and C are a.u.<sup>3</sup> mm s<sup>-1</sup>, mm s<sup>-1</sup>, and a.u.<sup>-3</sup>, respectively. R<sup>2</sup> is the coefficient of determination from the linear fit.

<sup>[</sup>c] Mean absolute deviations of the calculated ISs with respect to the experimental ISs, where the former ones are obtained by using Eq.(3).

<sup>[</sup>d]Maximum deviation of the calculated ISs from the experimental ISs.

<sup>[</sup>e]Standard deviation of the calculated ISs.

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Our evaluation included first some standard basis sets. With the default x2c-TZVPPall basis set le Online the calculated  $\rho(0)$  values, which are reflected in the very large value of C in Eq. (3), are significantly smaller than the fully relativistic electron density 15070 a.u.<sup>-3</sup>.86 The coefficient of determination (R<sup>2</sup>) obtained from linear regression analysis is very low and the value of "a" is far from the experimentally determined value (see Table 1). As a result, the MAD of the calculated isomer shifts from the experimental ones is also high (0.108 mm s<sup>-1</sup>). This is to be expected because of the lack of enough tight s basis functions in the x2c-TZVPPall basis set of Fe, which has been proven to be critical to obtain correct  $\rho(0)$  values.<sup>11</sup> For the same reason, shifting from the default x2c-TZVPPall to Weigend's segmented contracted relativistic basis set for NMR shielding constants (x2c-TZVPPall-s),<sup>62</sup> which has no additional s-space flexibility, does not result in any improvement.

The CP(PPP) basis set was originally proposed precisely for the prediction of Mössbauer parameters, albeit in a non-relativistic context.<sup>11</sup> Nevertheless, we test it and is a clear improvement also using the X2C Hamiltonian, underlining the leading importance of core flexibility for these properties. However, even though the calculated contact densities are better than those obtained with x2c-TZVPPall, those are still significantly smaller than 15070 a.u.<sup>-3</sup>. Next, we test the aug-cc-pVTZ-J basis set, which was optimized by Sauer and coworkers for the calculation of electron paramagnetic resonance (EPR) hyperfine coupling constants.<sup>63</sup> Interestingly, we get a good  $R^2$  value from the linear fit and the MAD is close to what was obtained with CP(PPP). However, the calculated  $\rho(0)$  values are very small and consequently the "a" value obtained from linear fit significantly deviates from the experimental calibration constant  $\alpha = -0.31 \pm 0.04$ . The probable reason behind this unusually small contact densities could be the presence of contracted s functions in aug-cc-pVTZ-J, which restricts the core flexibility of Fe. Due to the lack of sufficiently tight s-primitives, the  $\rho(0)$  values calculated using the relativistically contracted ANO-family basis set for iron, ANO-RCC-VTZP,66 has very poor linear correlation (R<sup>2</sup>=0.414) with the experimental isomer shifts. As a result, the fitted parameter "a" deviates significantly from the experimentally determined value (see Table 1).

Another standard basis set we have tested for Fe is DKH-def2-TZVPP, where the exponents of the Online def2-TZVPP64 were recontracted for the DKH2 Hamiltonian with a looser contraction. 43-47, 65 Although the calculated  $\rho(0)$  values are still not close to the fully relativistic density, the R<sup>2</sup> and "a" obtained from the linear regression analysis are considerably improved. Additionally, the mean absolute error and standard deviation of  $\delta_{cal}$  relative to  $\delta_{exp}$  are significantly better than those obtained with the x2c-TZVPPall basis set. The importance of flexibility in the s-functions is highlighted when we look at the correlation between the calculated  $\rho(0)$  and experimental isomer shifts with the s-decontracted x2c-TZVPPall, which is clearly better than standard x2c-TZVPPall. By using s-decontracted aug-cc-pVTZ-J the calculated contact densities of the iron centre are very close to the fully relativistic value 15070 a.u.<sup>-3</sup>. Moreover, we also achieve a very good correlation between  $\rho(0)$  and  $\delta_{exp}$  (R<sup>2</sup>=0.958).

Finally, we test two more modifications of aug-cc-pVTZ-J: (a) fully decontracted s functions and removal of the three innermost s-primitives (i.e., aug-cc-pVTZ-Jmod),<sup>67</sup> (b) fully decontracted s-functions and removal of the outermost d-, f-, and g-primitives (denoted sdecontracted aug-cc-pVTZ-J(-dfg)). The first one was recommended for the prediction of Cu(II) hyperfine coupling constants in a scalar relativistic approach,<sup>67</sup> whereas the second modification might be useful for large Fe-complexes where linear dependencies in the basis set may arise if diffuse functions are included. These modified basis sets yield effectively indistinguishable results. However, the calculated  $\rho(0)$  values with s-decontracted aug-ccpVTZ-J(-dfg) are closer to the fully relativistic electron density (i.e., 15070 a.u.<sup>-3</sup>) than those obtained with aug-cc-pVTZ-Jmod (see Table 1). With s-decontracted aug-cc-pVTZ-J and its two modifications, the mean absolute errors and the R<sup>2</sup> and "a" values obtained from the linear regression analysis are the same. We note that using a set of 12 iron clusters, Kurian and Filatov also found that the computed isomer shift values are only marginally affected by the addition (or subtraction) of the tightest primitive functions to a sizable, uncontracted basis set. 88 Unlike that study, however, here we find that the performance of contracted basis sets is noticeably worse compared to the large decontracted ones for the MPMIC80 set.

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For each basis set, the mean absolute error of the isomer shifts calculated using the point nucleus decention model is slightly higher than those obtained with the Gaussian finite nucleus model (see Table 1 and Table S4 in the ESI). Except for the aug-cc-pVTZ-J and its modifications, the linear fitting parameters of other basis sets are quite similar regardless of whether a point nucleus or finite nucleus model is used.

Our conclusion regarding basis set selection for the calculation of the Mössbauer isomer shifts is that within the X2C approach with the Gaussian finite nucleus model a basis set with very tight *s* functions is necessary. Among the standard and non-standard basis sets evaluated here, *s*-decontracted versions of aug-cc-pVTZ-J and aug-cc-pVTZ-J(-*dfg*) are equally good and are among the best choices. For the next step, we benchmark different DFT methods in combination with *s*-decontracted aug-cc-pVTZ-J, CP(PPP), and DKH-def2-TZVPP.

## 4.1.2. Comparison of DFT methods

For each of the three selected basis sets, fifteen different density functional approximations are tested. The parameters obtained from the linear regression analysis, coefficient of determination (R<sup>2</sup>), and error statistics of  $\delta_{cal}$  with respect to  $\delta_{exp}$  are listed in Table 2. For the calibration parameters and error statistics obtained using the *s*-decontracted aug-cc-pVTZ-J(-*dfg*) basis set for Fe, see Table S5 in ESI.

Unlike the different basis sets, using different density functionals does not have much influence on the calculated contact densities. Based on the R<sup>2</sup> and mean absolute errors, climbing the rungs of "Jacob's ladder" improves accuracy gradually from the 1<sup>st</sup> to the 4<sup>th</sup> rung. As in earlier findings, hybrid functionals demonstrate a significant improvement over the performance of pure GGA and meta-GGA approaches.<sup>26, 27, 33, 88, 89</sup> The inferior correlations obtained with pure density functionals can be attributed to their incorrect behavior near the nucleus.<sup>90</sup> The only exception is the extensively parametrized hybrid functional M06, which offers accuracy similar to the pure meta-GGA functionals. Among the hybrid functionals, range separation offers no benefit over the global hybrid variants, and it does more harm than good when there is no short-range HF-exchange involved (e.g., in LC-BLYP). Contrary to what Römelt *et al.*<sup>18</sup> found with the scalar-relativistic ZORA Hamiltonian<sup>41,91,92</sup> on a much smaller dataset, the performance of

the double hybrid functional for the extensive MPMIC80 set is inferior to standard hybrids in

**Table 2**: Calibration constants and statistical parameters from the linear regression analysis of the computed contact densities using different DFT functionals and the experimental isomer shifts.<sup>[a]</sup>

SVWNS   -0.32   0.8946   14831   0.913   0.071   0.296   0.00	Basis set for Fe	Methods	а	b	C	R <sup>2</sup>	MAD <sup>[b]</sup> (mm s <sup>-1</sup> )	Max. Dev. <sup>[c]</sup> (mm s <sup>-1</sup> )	St. Dev. <sup>[d]</sup> (mm s <sup>-1</sup> )
BB86		SVWN5	-0.32	0.8946	14831	0.913		<del></del>	0.098
PBE   -0.32   0.9970   14933   0.921   0.067   0.267   0.0		BP86					0.067		0.093
BLYP				0.9970	14933		0.067		0.094
s-decontracted aug-cc-pVTZ-J  BBLYP		BLYP	-0.32	1.0974	14954	0.917	0.070		0.096
Sedecontracted aug-cc-pVTZ-J		TPSS		1.0045	14917	0.935	0.060	0.243	0.085
Balyp		PBE0	-0.28	1.1225	14920	0.965	0.050	0.148	0.063
Bally   -0.29   1.1073   14930   0.958   0.053   0.180   0.0   0.0000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.00000   0.000000   0.000000   0.000000   0.000000   0.000000   0.000000   0.0000000   0.00000000	1 , , 1								0.066
TPSS0							0.053		0.069
TPSS0	aug-cc-pv1Z-J	TPSSh	-0.30	1.1976	14913	0.954	0.052	0.227	0.072
M06			1					0.122	0.060
LC-BLYP				0.9446	15010			0.277	0.084
CAM-B3LYP		LC-BLYP	-0.29	1.0845	14951	0.950	0.062	0.168	0.075
									0.067
B2PLYP									0.068
SVWN5									0.083
BP86									0.100
PBE									0.095
BLYP									0.096
TPSS									0.098
PBEO									0.087
CP(PPP)    B3LYP									0.063
CP(PPP)  B3LYP									0.064
TPSSh	CP(PPP)								0.070
TPSS0	01(111)		1						0.073
M06									0.062
LC-BLYP									0.085
CAM-B3LYP ωB97X         -0.29         1.0531         14370         0.961         0.054         0.134         0.0           B2PLYP         -0.28         1.1687         14450         0.959         0.055         0.147         0.0           B2PLYP         -0.25         1.1856         14350         0.940         0.066         0.176         0.0           SVWN5         -0.33         0.9272         14010         0.909         0.074         0.305         0.1           BP86         -0.33         0.9640         14094         0.919         0.069         0.277         0.0           PBE         -0.33         1.1343         14086         0.919         0.069         0.273         0.0           BLYP         -0.33         1.0651         14089         0.914         0.072         0.280         0.0           TPSS         -0.33         1.1151         14078         0.934         0.061         0.247         0.0           DKH-def2-         PBE0         -0.29         1.0853         14078         0.964         0.051         0.156         0.0           B1LYP         -0.29         1.1600         14080         0.962         0.054         0.161         0.0									0.074
B97X B2PLYP         −0.28 −0.25         1.1687 1.1856         14450 14350         0.959 0.940         0.055 0.066         0.147 0.066         0.0           SVWN5         −0.33         0.9272         14010         0.909         0.074         0.305         0.1           BP86         −0.33         0.9640         14094         0.919         0.069         0.277         0.0           PBE         −0.33         1.1343         14086         0.919         0.069         0.273         0.0           BLYP         −0.33         1.0651         14089         0.914         0.072         0.280         0.0           TPSS         −0.33         1.1151         14078         0.934         0.061         0.247         0.0           PBE0         −0.29         1.0853         14078         0.944         0.051         0.156         0.0           B1LYP         −0.29         1.1600         14080         0.962         0.054         0.161         0.0           TZVPP         B3LYP         −0.30         1.0315         14076         0.956         0.055         0.190         0.0           TPSS0         −0.29         1.0877         14072         0.967         0.049         0.127									0.066
B2PLYP         -0.25         1.1856         14350         0.940         0.066         0.176         0.0           SVWN5         -0.33         0.9272         14010         0.909         0.074         0.305         0.1           BP86         -0.33         0.9640         14094         0.919         0.069         0.277         0.0           PBE         -0.33         1.1343         14086         0.919         0.069         0.273         0.0           BLYP         -0.33         1.0651         14089         0.914         0.072         0.280         0.0           TPSS         -0.33         1.1151         14078         0.934         0.061         0.247         0.0           PBE0         -0.29         1.0853         14078         0.964         0.051         0.156         0.0           PBE0         -0.29         1.1600         14080         0.962         0.054         0.161         0.0           B1LYP         -0.30         1.0315         14076         0.956         0.055         0.190         0.0           TPSSh         -0.32         0.9873         14076         0.953         0.053         0.220         0.0           TPSSO <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>0.068</td>									0.068
SVWN5         -0.33         0.9272         14010         0.909         0.074         0.305         0.1           BP86         -0.33         0.9640         14094         0.919         0.069         0.277         0.0           PBE         -0.33         1.1343         14086         0.919         0.069         0.273         0.0           BLYP         -0.33         1.0651         14089         0.914         0.072         0.280         0.0           TPSS         -0.33         1.1151         14078         0.934         0.061         0.247         0.0           PBE0         -0.29         1.0853         14078         0.964         0.051         0.156         0.0           PBE0         -0.29         1.1600         14080         0.962         0.054         0.161         0.0           B1LYP         -0.30         1.0315         14076         0.956         0.055         0.190         0.0           TPSSh         -0.32         0.9873         14076         0.953         0.053         0.220         0.0           TPSSO         -0.29         1.0877         14072         0.967         0.049         0.127         0.0           M06									0.082
BP86									0.101
DKH-def2- TZVPP    PBE									0.095
DKH-def2- TZVPP    BLYP									0.095
DKH-def2- TZVPP  TPSS									0.098
DKH-def2- TZVPP    PBE0			1						0.086
DKH-def2- TZVPP         B1LYP B3LYP         -0.29 -0.30         1.1600 1.0315         14076 14076         0.962 0.956         0.054 0.055         0.161 0.190         0.0 0.0           TPSSh TPSS0         -0.32 -0.29         0.9873 1.0877         14076 14072         0.967 0.967         0.049 0.049         0.127 0.049         0.0 0.282         0.0 0.0 0.067         0.282 0.0 0.065         0.065 0.163         0.0 0.065         0.163 0.0 0.065         0.0 0.082         0.0 0.0 0.082         0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0			1						0.063
DKH-det2- TZVPP         B3LYP TPSSh         -0.30         1.0315         14076         0.956         0.055         0.190         0.0           TPSSh         -0.32         0.9873         14076         0.953         0.053         0.220         0.0           TPSS0         -0.29         1.0877         14072         0.967         0.049         0.127         0.0           M06         -0.32         1.2105         14105         0.934         0.067         0.282         0.0           LC-BLYP         -0.30         1.0128         14087         0.945         0.065         0.163         0.0           CAM-B3LYP         -0.29         1.2614         14081         0.957         0.058         0.138         0.0									0.065
TPSSh			1						0.070
TPSS0     -0.29     1.0877     14072     0.967     0.049     0.127     0.0       M06     -0.32     1.2105     14105     0.934     0.067     0.282     0.0       LC-BLYP     -0.30     1.0128     14087     0.945     0.065     0.163     0.0       CAM-B3LYP     -0.29     1.2614     14081     0.957     0.058     0.138     0.0			1						0.070
M06       -0.32       1.2105       14105       0.934       0.067       0.282       0.0         LC-BLYP       -0.30       1.0128       14087       0.945       0.065       0.163       0.0         CAM-B3LYP       -0.29       1.2614       14081       0.957       0.058       0.138       0.0									0.072
LC-BLYP     -0.30     1.0128     14087     0.945     0.065     0.163     0.0       CAM-B3LYP     -0.29     1.2614     14081     0.957     0.058     0.138     0.0									0.086
CAM-B3LYP   -0.29									0.080
									0.078
$\begin{bmatrix} \omega_{D7/A} & \end{bmatrix} = 0.26 + 1.2300 + 14007 + 0.937 & 0.030 + 0.140 & 0.0$			1						0.069
B2PLYP -0.25 1.1934 14070 0.938 0.66 0.178 0.0									0.069

 $<sup>^{[</sup>a]}a$  and b are the fitting coefficients obtained from the linear fit of Eq. (3). C is a constant, which is very close to the calculated  $\rho(0)$  value. The units of a, b, and C are a.u.<sup>3</sup> mm s<sup>-1</sup>, mm s<sup>-1</sup>, and a.u.<sup>-3</sup>, respectively. R<sup>2</sup> is the coefficient of determination from the linear fit.

<sup>[</sup>b] Mean absolute deviations of the calculated ISs with respect to the experimental ISs, where the former ones are obtained by using Eq.(3).

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For the TPSS exchange- and correlation-based hybrid functionals, increasing the percentage of exact exchange from 10% to 25% improves its accuracy. Therefore the question may arise, what is the optimum percentage of exact exchange for predicting the isomer shifts in MPMIC80? To investigate this, we chose the BLYP and TPSS functionals and their hybrid counterparts with varying percentage of exact (Hartree–Fock, HF) exchange (i.e., %HFx) while employing the s-decontracted aug-cc-pVTZ-J basis set for iron. Polynomial fitting of MADs with respect to the %HFx suggest minima near 30% for the BLYP-based and near 25% for the TPSS-based functionals (see Figure 3 and Table S6 in the ESI). The MAD gap between the BLYP- and TPSS-based functionals decreases with the increase of exact exchange. We obtained the lowest mean absolute error with TPSS0 (25% HFx) among all functionals, and it starts to increase rapidly with higher percentage of exact exchange. For the present dataset the fitting parameter "a" from linear regression has strong dependence on the exact exchange, but it is not sensitive to specific exchange and correlation combinations (Figure S1). Finally, we note that despite having 27% HF exchange, the M06 functional gives inferior electron densities and hence its performance for isomer shifts is inferior to other hybrid functionals (Table 2).

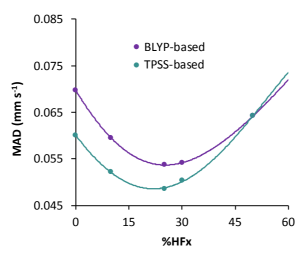


Figure 3: Effect of the percentage of exact exchange (%HFx) used in BLYP- and TPSS-based hybrid functionals on the MAD values of the calculated isomer shifts relative to the experimentally measured shifts.

As the GGA exchange and correlation parts of B2PLYP (i.e., B88 and LYP) work fine for control dataset, the origin of the poor performance of the double hybrid functional could be either the PT2 correlation part or the amount of exact exchange used. We note that for the calculation of Cu(II) hyperfine coupling constants, the use of relaxed densities for the PT2 part was recommended.<sup>67</sup> Following the same protocol and using the *s*-decontracted aug-cc-pVTZ-J basis set for Fe improves the "*a*" value of the linear fit from –0.24 to –0.27. However, R² and mean absolute error are worse than the unrelaxed density calculations, allowing us to rule out the first possible source of error (see Table S7). Hence, the high fraction of exact exchange (53%) used in the B2PLYP seems to be the principal reason behind its poorer performance compared to standard hybrid DFT functionals.

Earlier studies have shown that spin-component scaled double hybrids are more accurate than simple double hybrids like B2PLYP for calculating the energetics and spectral properties of transition metal complexes. 93-96 The contact densities computed using the DSD-PBEP86 functional and the *s*-decontracted aug-cc-pVTZ-J basis set for iron, along with the relaxed PT2 density, were fitted to the experimental isomer shifts. This resulted in the fitting parameter a = -0.26 with  $R^2 = 0.895$  (see Figure S2). The relatively high percentage (68%) of HF exchange used in DSD-PBEP86 could be the reason behind its marginally poorer performance than B2PLYP. However, for most of the quintet Fe<sup>II</sup> and sextet Fe<sup>III</sup> complexes of our dataset, the isomer shifts calculated using DSD-PBEP86 are closer to the experimental values than  $\delta_{cal}$  using TPSS0 or B2PLYP (see Table S11). We note in passing that a double hybrid calculation with relaxed density is approximately three times more expensive and requires more memory allocation than a similar calculation with unrelaxed density. The "unrelaxed" PT2 density corresponds to simple PT2 expectation value density, whereas the "relaxed" one incorporates orbital relaxation.

In conclusion, hybrid functionals incorporating ca. 25–30% exact exchange offer the best linear correlation and accuracy for predicting isomer shifts. Among all methods tested, our best picks are TPSS0 and PBE0, combined with the *s*-decontracted aug-cc-pVTZ-J basis set for Fe.

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With respect to the literature, we note that using 15 iron-containing compounds and explosive Online CP(PPP) basis set for Fe, an R<sup>2</sup>=0.972 was obtained from the linear regression analysis of the nonrelativistic electron densities calculated with B3LYP against experimental isomer shifts.<sup>11</sup> The reported standard deviation for the theoretical prediction of isomer shifts was 0.09 mm s<sup>-1</sup>. Later, with a slightly larger and more diverse set, B3LYP offered a marginally better correlation (R<sup>2</sup>=0.980) and a standard deviation of 0.09 mm s<sup>-1</sup>, while using a larger dataset, Pápai and Vankó found R<sup>2</sup>=0.975 and MAD=0.06 mm s<sup>-1</sup> for B3LYP.<sup>30</sup> However, Kurian and Filatov found a better correlation with experimental results using BH&HLYP instead of B3LYP, which might be a result of employing a small dataset. 88 Using a dataset comprising 69 iron compounds, Comas-Vilà and Salvador showed that replacing the density at the iron nucleus with the density integrated in a sphere of radius 0.06 au surrounding the iron center can provide excellent correlation ( $R^2 = 0.976$ ) when using the conventional def2-TZVP basis set.<sup>89</sup> In a recent study, using 20 molecular Fe complexes, Gallenkamp et al. found the best performance with TPSSh and PBE0 (R<sup>2</sup>= 0.978 and 0.976) with mean absolute errors of 0.05 and 0.06 mm s<sup>-1</sup>, respectively.<sup>33</sup> With a set of 21 iron-complexes, using the scalar-relativistic ZORA Hamiltonian and ZORA-def2-TZVP basis set for Fe, hybrid, and double hybrid functionals offered similar accuracy (R<sup>2</sup>=0.970 and Standard deviation=0.08 mm s<sup>-1</sup>). <sup>18</sup> Employing the DKH2 Hamiltonian and custom-defined basis sets for Fe did not offer any further improvement in the performance of hybrid functionals.<sup>22</sup> Although our best pick, TPSS0 with the s-decontracted aug-cc-pVTZ-J basis set for Fe, has a slightly smaller R<sup>2</sup> value, the standard deviation is better than what was reported previously. The smaller R<sup>2</sup> can be attributed to using a dataset that is more than three times larger than what was used in refs. 18, 22 Perdew and coworkers argue that highly parameterized density functional methods are often significantly inferior to the functionals developed by constraint satisfaction while calculating the electron densities.<sup>97</sup> Among the 128 functionals tested in their work, PBE0 is one of the best for calculating electron density distributions compared to the all-electron coupled cluster singles and doubles (CCSD-full) densities. PBE0 and TPSS0 should therefore yield reliable electron densities and, hence, good performance for Mössbauer isomer shift computations.

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## 4.2. High-Temperature Isomer Shifts

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It is well known that the Mössbauer parameters are temperature-dependent, and this behavior originates from the temperature dependence of the Lamb–Mössbauer factor f. With increasing temperature, the second-order Doppler shift (SODS) appears owing to significant thermal motions of the source and absorber nuclei, which leads to a lower isomer shift.<sup>5, 9</sup> The SODS-corrected chemical isomer shift of a Mössbauer spectrum is sometimes referred to in literature as the Center Shift (CS),<sup>98</sup> which has the following expression

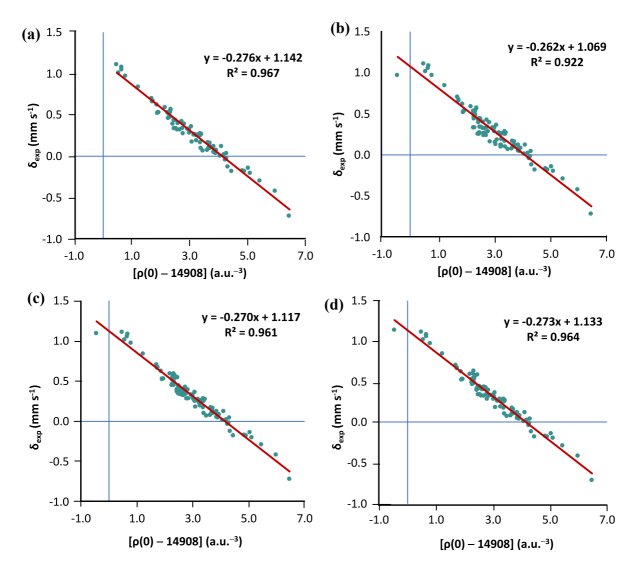
$$CS = \delta + SODS \tag{5}$$

Although at temperature up to 77 K the SODS contributes less than 0.02 mm s<sup>-1</sup> to the measured isomer shift, it can be as large as 0.1 mm s<sup>-1</sup> at room temperature. A fair comparison between computational and experimental isomer shifts is therefore guaranteed only at temperatures of a few Kelvin (K). Since the influence of the temperature may exceed the uncertainties from density functional methods and basis sets, comparison with experimental data obtained above 80 K needs special treatment before comparing them with the calculated counterparts. In multiple occasions, Noodleman and co-workers have proposed a linear SODS correction of the experimentally recorded isomer shifts by 0.12 mm s<sup>-1</sup> for a temperature decrease from 300 K to 4.2 K.<sup>25, 99-101</sup> However, Friesner and co-workers recommended to avoid such empirical corrections in a benchmark study because of the nontrivial temperature dependence of the <sup>57</sup>Fe Mössbauer parameters.<sup>27</sup>

As we already have a curated subset of molecules (21 Fe-complexes) with high-T experimental isomer shirts, we can examine whether the solution proposed by Noodleman is transferable (see Table S2 in the ESI). For this purpose, we chose the TPSS0 functional with the s-decontracted aug-cc-pVTZ-J basis set for iron. Considering the 80 low-temperature isomer shifts, linear regression analysis of  $\rho(0)$  vs.  $\delta_{\rm exp}$  gives R<sup>2</sup>=0.967. Adding the 21 high-T isomer shifts to the MPMIC80 and refitting the linear equation we obtain a significantly lower R<sup>2</sup> value of 0.922. However, if Noodleman's correction is applied to the 21 high-T isomer shifts, linear regression analysis for the mixed set of 101 compounds almost recovers the correlation obtained with the

original MPMIC80 set (see Figure 4a-c). The same trend is observed with PBE0/s-decontracted lete Online aug-cc-pVTZ-J(Fe) and B3LYP/CP(PPP)(Fe) (see Figures S3 and S4).

Now, refitting a high-T isomer shift correction factor against 21 complexes we get the value 0.16 mm s<sup>-1</sup>. However, using a larger IS correction factor only marginally improves correlation compared to Noodleman's correction (see Figure 4c-d).



**Figure 4:** Plot of the calculated TPSS0/s-decontracted aug-cc-pVTZ-J+x2c-TZVPPall level contact densities against the (a) low-T isomer shifts of MPMIC80; (b) 80 low-T and 21 high-T isomer shifts; (c) 80 low-T isomer shifts and 21 isomer shifts with Noodleman's high-T SODS correction (0.12 mm s<sup>-1</sup>) and (d) 80 low-T isomer shifts and 21 isomer shifts with refitted high-T SODS correction (0.16 mm s<sup>-1</sup>).

## 4.2. Quadrupole Splittings in the MPMIC80 Set

View Article Online DOI: 10.1039/D4CP00431K

Similar to isomer shifts, we start with the non-relativistic quadrupole splitting calculation using TPSSh functional and CP(PPP) basis set for Fe. Relative to the experimental  $|\Delta E_Q|$ , non-relativistically obtained quadruple splittings yield mean absolute deviation and root-mean-square deviation 0.331 and 0.443 mm s<sup>-1</sup>, respectively. Although deviation from  $|\Delta E_{Q,exp}|$  is distributed among all the oxidation states of Fe, the largest error was obtained for iron(III) phthalocyanine chloride and octaethylporphyrinato-iron(II) (i.e., molecules #35 and #65 from the MPMIC80 dataset; see Figure S5).

**Table 3:** For ten basis sets, the mean absolute deviations (MADs) and root-mean-square deviations (RMSDs) of calculated quadrupole splittings with respect to the experimental values. The scalar-relativistic calculations were performed with B3LYP functional and finite nucleus model.

Basis set for Fe	MAD(mm s <sup>-1</sup> )	RMSD(mm s <sup>-1</sup> )
CP(PPP)	0.312	0.451
x2c-TZVPPall	0.357	0.504
x2c-TZVPPall-s	0.299	0.436
aug-cc-pVTZ-J	0.352	0.495
DKH-def2-TZVPP	0.241	0.381
ANO-RCC-VTZP	0.283	0.421
s-decontracted x2c-TZVPPall	0.361	0.514
s-decontracted aug-cc-pVTZ-J	0.347	0.487
s-decontracted aug-cc-pVTZ-J (-dfg)	0.349	0.489
aug-cc-pVTZ-Jmod <sup>67</sup>	0.347	0.487
Non-Relativistic <sup>[b]</sup>	0.331	0.443

<sup>[</sup>a] For the results with point nucleus model, see Table S8 in the supporting information.

[b] with CP(PPP) basis set for Fe and def2-TZVP for the ligand atoms.

The mean absolute and root-mean-square errors of calculated  $|\Delta E_Q|$  with respect to the experimental  $|\Delta E_Q|$  for ten different basis sets evaluated using B3LYP functional, finite nucleus model, and first-order picture change effect are listed in Table 3. Among the six standard basis sets, DKH-def2-TZVPP and x2c-TZVPPall are the best and the worst performer, respectively. Unlike what we observed for isomer shifts, x2c-TZVPPall-s basis set, developed for NMR shielding constants, offers a noticeably better accuracy than x2c-TZVPPall. Decontraction of the *s* primitives does more harm than good for x2c-TZVPPall.

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Table 4: Performance statistics of fourteen different density functionals for the calculation of Tester Online quadrupole splittings in MPMIC80 dataset.

Basis set for Fe	Methods	MAD (mm s <sup>-1</sup> )	RMSD (mm s <sup>-1</sup> )	
	SVWN5	0.319	0.495	
	BP86	0.274	0.450	
	PBE	0.280	0.457	
	BLYP	0.272	0.450	
	TPSS	0.269	0.437	
- decembrants d	PBE0	0.374	0.523	
	B1LYP	0.404	0.557	
s-decontracted	B3LYP	0.347	0.487	
aug-cc-pVTZ-J	TPSSh	0.261	0.410	
	TPSS0	0.383	0.525	
	M06	0.544	0.949	
	LC-BLYP	0.432	0.562	
	CAM-B3LYP	0.474	0.632	
	ωB97X	0.493	0.649	
	B2PLYP	0.648	0.857	
	SVWN5	0.348	0.516	
	BP86	0.308	0.473	
	PBE	0.319	0.484	
	BLYP	0.303	0.475	
	TPSS	0.305	0.463	
	PBE0	0.326	0.457	
	B1LYP	0.346	0.480	
CP(PPP)	B3LYP	0.312	0.451	
CI (III)	TPSSh	0.251	0.405	
	TPSS0	0.353	0.489	
	M06	0.500	0.891	
	LC-BLYP	0.371	0.487	
	CAM-B3LYP	0.409	0.553	
	ωΒ97Χ	0.419	0.546	
	B2PLYP	0.587	0.779	
	SVWN5	0.451	0.602	
	BP86	0.398	0.545	
	PBE	0.408	0.557	
	BLYP	0.396	0.542	
	TPSS	0.396	0.534	
	PBE0	0.258	0.399	
DKH-def2-TZVPP	B1LYP	0.258	0.392	
	B3LYP	0.238	0.381	
	TPSSh	0.291	0.431	
	TPSS0	0.291	0.412	
	M06	0.270	0.412	
	LC-BLYP	0.423	0.813	
	CAM-B3LYP	0.301	0.440	
	ωB97X	0.305	0.423	
	B2PLYP	0.466	0.635	

Next, for the evaluation of DFT methods to calculate  $|\Delta E_Q|$  values three basis sets were chosen: CP(PPP), DKH-def2-TZVPP, and s-decontracted aug-cc-pVTZ-J. The MAD and RMSD statistics for each the method and basis set combinations are listed in Table 4. (For the error

statistics obtained using the *s*-decontracted aug-cc-pVTZ-J(-dfg) basis set for Fe, see Table SQCIe Online in the supporting information.)

For all these basis sets, GGA and *meta*-GGA functionals offer better accuracy than the LDA functional. Importantly, and unlike what we observed for the isomer shifts, hybrid functionals produce larger errors than GGA and *meta*-GGA functionals. Range separation leads to deterioration rather than improvement. The only exception is when the DKH-def2-TZVPP functional is used, in which case hybrid functionals are better performers than the 2<sup>nd</sup> and 3<sup>rd</sup> rung functionals. Interestingly, using either the CP(PPP) or the *s*-decontracted aug-cc-pVTZ-J basis set for Fe combined with TPSSh functional provides noticeably lower mean absolute error than TPSS0.

Once again, B2PLYP is the worst performer among all functionals tested. While using the s-decontracted aug-cc-pVTZ-J basis set for Fe, employing relaxed density for PT2 correlation improve its accuracy (MAD goes down from 0.648 to 0.485 mm s<sup>-1</sup>). A large percentage of exact exchange in B2PLYP also contributes a significant share to its large mean absolute error. Analyzing the results obtained from varying percentage of exact exchange in PBE-, BLYP-, and TPSS-based hybrid functionals, it is evident that beyond 10% the mean absolute error in calculated  $|\Delta E_Q|$  compared to experiment increases rapidly (see Table S10 in the supporting information). However, pure GGA and *meta*-GGA functionals are also good and cheaper alternatives. This observation agrees with the recommendation of Nemykin and Hadt for using the pure GGA functional BPW91 over B3LYP to predict quadrupole splittings in ferrocenes accurately.<sup>26</sup> Overall, the quadrupole splittings are harder to reproduce systematically with equally high level of reliability as the isomer shifts and remain a challenge that will have to be addressed more satisfactorily in future studies.

So far, we have used the  $^{57}$ Fe nuclear quadrupole moment Q = 0.16 b to calculate the quadrupole splitting values. A reasonable question is whether there is anything to be gained by refitting the quadrupole moment of the  $^{57}$ Fe nucleus against the MPMIC80 set. Using the s-decontracted aug-cc-pVTZ-J basis set for iron, we selected nine representative functionals for this purpose and plotted the experimental QSs as a function of

$$1/2eV_{zz}\sqrt{1+\frac{1}{3}\left(\frac{|V_{yy}-V_{xx}|}{V_{xx}}\right)^2}$$
 (5) View Article Online DOI: 10.1039/D4CP00431K

Following Eq. (4), the new  $Q(^{57}Fe)$  value is determined from the slope of the linear fit to the plot (see Table 5). With TPSS-based hybrid functionals, increasing the percentage of HF-exchange gradually decreases the fitted  $Q(^{57}Fe)$ .

Using the nuclear quadrupole moment thus obtained, we reevaluated the quadrupole splittings and calculated the MAD and RMSD errors against the experimental  $\Delta E_Q$  values (see Table 5). It can be seen that refitting the <sup>57</sup>Fe NQMs only marginally improves the performance of pure GGA and mGGA functionals. This improvement is negligible for the hybrid functionals at a small fraction of HF exchange, and increasing the %HFx also increases the performance gap.

**Table 5:** Re-determined values for the iron nuclear quadrupole moment  $Q(^{57}Fe)$  and the error statistics obtained with various methods against the experimental quadrupole splittings. The *s*-decontracted aug-cc-pVTZ-J basis set was employed for Fe.

Functionals	Calculated $Q(^{57}\mathrm{Fe})^{[a]}$	MAD (mm s <sup>-1</sup> )	RMSD (mm s <sup>-1</sup> )
BP86	0.165	0.268	0.446
PBE	0.166	0.271	0.452
TPSS	0.167	0.254	0.429
TPSSh	0.155	0.248	0.405
B3LYP	0.141	0.272	0.408
PBE0	0.139	0.285	0.424
TPSS0	0.141	0.303	0.435
TPSS30	0.137	0.328	0.456
TPSS50	0.125	0.429	0.569

[a] slope of the linear fit to the plot of experimental QS vs.  $\frac{1}{2}eV_{zz}\sqrt{1+\left(|V_{yy}-V_{xx}|/V_{zz}\right)^2/3}$ 

## 5. Conclusions

We have presented the largest and most extensive molecular dataset, namely MPMIC80, for the calculation of <sup>57</sup>Fe Mössbauer isomer shifts and quadruple splittings. This new set of complexes covers most of the known spin and oxidation states of iron, a diverse set of ligands, and with each complex having one unique iron center and low-temperature experimentally determined Mössbauer parameters. After an extensive survey of density functional methods and basis sets with the aid of the MPMIC80 data set, we can conclude the following:

- Irrespective of the choice of basis sets and density functional methods, use of first clear change effect is recommended with the scalar relativistic X2C-Hamiltonian.
- Tight s-primitives are necessary for the calculation of contact densities close to the fully relativistic limit ( $\rho = 15070 \text{ a.u.}^{-3}$ ). Consequently, standard basis sets that are either general-purpose or have been optimized for other properties (e.g., for NMR chemical shifts) are inappropriate poor choices for these calculations, whereas the s-decontracted version of aug-cc-pVTZ-J is a solid choice.
- For isomer shifts, 25-30% is the optimum percentage of exact (HF) exchange for hybrid functionals with scalar relativistic X2C Hamiltonian. Best accuracy was found with the TPSSO and PBEO functionals with s-decontracted aug-cc-pVTZ-J. Range separation has no benefit over global hybrid functionals.
- Double hybrid functionals are not recommended. The performance of B2PLYP is worse
  than any hybrid functional, which is most likely the result of the high percentage of
  exact exchange used.
- Refitting the original 80 isomer shifts and 21 high-T isomer shifts adjusted to 4.2 K by a linear correction against the calculated contact densities almost fully recovers the  $R^2$  value obtained with the low-temperature MPMIC80.
- Quadruple splitting values are less systematically predicted with DFT. Unlike isomer shifts, pure GGA, meta-GGA, and hybrid functionals with small fraction of HF exchange (~10%) are preferred for these calculations, but there is considerable room for improvement.

## Acknowledgment

The authors gratefully acknowledge support from the Max Planck Society. G.S. thanks Dr. Daniel SantaLucia (Max Planck Institute for Chemical Energy Conversion) and Dr. Georgi Stoychev (FAccTs GmbH, Germany) for helpful discussions. G.S. and D.A.P. thank Dr. Sergio A. V. Jannuzzi (Max Planck Institute for Chemical Energy Conversion) for pointing out a recently reported Fe<sup>VII</sup> complex.

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Additional information

View Article Online DOI: 10.1039/D4CP00431K

Supplementary information is available in the online version of the paper.

## **Competing financial interests**

The authors declare no competing financial interests.

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View Article Online DOI: 10.1039/D4CP00431K

## Data availability statement

The data supporting this article have been included as part of the Supplementary Information.