Theoretical investigation on the presence of different iron sites in RbMn[Fe(CN)₆]·H₂O

Reshmi Kurian and Michael Filatov
Theoretical Chemistry, Zernike Institute for Advanced Materials, Rijksuniversiteit Groningen, Nijenborgh 4, 9747 AG Groningen, The Netherlands
E-mail: m.filatov@rug.nl

Abstract. The ⁵⁷Fe electron contact densities and the corresponding Mössbauer isomer shifts of RbMn[Fe(CN)₆]·H₂O are calculated with the use of ab initio wavefunction methods and double hybrid density functional method. The theoretical analysis of possible origin of the two slightly different signals in the Mössbauer spectra observed at 293 K and 50 K (Vertelman et al. 2008 Chem. Mater. 20 1236-38) is carried out. The measured differences of 0.083 - 0.105 mm/s between the high-temperature and low-temperature phases can be attributed to the different oxidation states of iron: the iron is in Fe³⁺ oxidation state in the high temperature phase and Fe²⁺ state in the low temperature phase. The smaller isomer shift differences of 0.012 - 0.034 mm/s occurring in both high temperature and low temperature structures can be attributed to different distributions of the Rb⁺ ions within the unit cell of RbMn[Fe(CN)₆]·H₂O.

1. Introduction
In the present work, theoretical analysis of ⁵⁷Fe Mössbauer isomer shifts in a Prussian blue analogue RbMn[Fe(CN)₆]·H₂O is carried out. Prussian blue analogues are compounds with the general molecular formula AₓMa[Mb(CN)₆]·zH₂O (where, A = alkali cation and Ma/Mb = metal ion), which attract considerable interest due to their peculiar magneto-optical properties [1]. Thus, RbMn[Fe(CN)₆]·H₂O shows light and temperature induced switching of magnetization [2]. Upon cooling from 300K to approximately 150K, RbMn[Fe(CN)₆]·H₂O undergoes transition to a state with higher molar magnetization. A reversed transition to a low magnetization state is observed upon heating to ca. 250K. It has been hypothesized that the switching of magnetization occurs due to electron transfer from the Mn²⁺ cations to the Fe³⁺ ions (and reversed transition) as a result of structural phase transition upon cooling/heating the samples [3].

The Mössbauer spectra of the high-temperature (HT) and the low-temperature (LT) structures were recorded whereby it was observed that, in both structures, two signals with the isomer shifts reported in Table 1 were present [3]. The presence of two different Fe³⁺ sites in the HT phase of

| T, K  | δ, mm/s       |
|-------|---------------|
| 50    | -0.033(4)     |
| 293   | -0.138(2)     |

Table 1. ⁵⁷Fe Mössbauer isomer shifts observed in RbMn[Fe(CN)₆]·H₂O at different temperatures [3] where number in the parentheses shows the experimental uncertainty in the last digit.
RbMn[Fe(CN)₆]·H₂O with the shifts \( \delta = -0.138(2) \) mm/s and \( \delta = -0.150(2) \) mm/s was interpreted as a consequence of different distributions of the Rb cations over the interstitial sites 4c and 4d (see Figure 1) in the space group \( \overline{F}43m \) of the crystal. Although it was not possible to obtain reliable crystal structures of the LT phase of RbMn[Fe(CN)₆]·H₂O, it is clear that iron is present in the form of low-spin Fe\(^{II}\) ions octahedrally coordinated by the cyano ligands. Similar to the HT phase, a disorder in the occupation of the Rb sites around the Fe\(^{II}\) ions leads to emergence of two different iron sites as can be seen from the isomer shifts reported in Table 1.

To obtain insight into the origin of different iron sites in the HT and LT structures of RbMn[Fe(CN)₆]·H₂O we undertake quantum chemical calculations of \(^{57}\)Fe Mössbauer isomer shifts in cluster models of these sites. More specifically, our goal is to elucidate whether the observed isomer shifts originate from variations in the oxidation state of iron ions (Fe\(^{II}\) vs. Fe\(^{III}\) isomer shifts) and from differences in the coordination sphere of iron ions due to different distribution of the Rb\(^{I}\) ions (see Figure 1).

### 2. Details of calculations

The calculations are carried out in the embedded cluster approach, where the geometries of clusters shown in Fig. 1 are obtained from crystal structure of the HT phase reported in Ref. [3]. The iron is in octahedral coordination with six cyano ligands bridging Fe and Mn. The disorder in the distribution of Rb\(^{I}\) ions is modeled by selecting two possible distribution patterns of Rb\(^{I}\) around iron atoms as shown in Fig. 1. The Fe-C and Mn-N distances are 1.929 Å and 2.205 Å respectively. The same geometry of clusters is used for the LT phase calculations. The Madelung field is modeled by the field of a large array of point charges placed at the appropriate crystallographic positions.

The calculations are carried out using Hartree-Fock (HF), second order Møller-Plesset perturbation theory (MP2) [4], Coupled Cluster Singles and Doubles (CCSD) [4] and double hybrid density functional (B2-PLYP [5]) level. The 24s12p9d basis set of Partridge with a set of polarization functions taken from TZVpp basis set of Ahlrichs and May [6] is used for iron and 6-31G* [7] basis set is used for carbon and nitrogen. All basis sets were used in uncontracted form. The Rb\(^{I}\) core is modeled by the Stuttgart effective core potential (ECP) [8], whereas Mn\(^{II}/^{III}\) cores are modeled by the representative ECPs of Ca\(^{II}\) and Al\(^{III}\), respectively. The closed shell systems are treated with spin restricted formalism and the open-shell systems with spin unrestricted formalism. The relativistic calculations are carried out within the one-electron approximation [9] and using the normalized elimination of the small component (NESC) [10] method which was implemented according to Ref. [11].

The \(^{57}\)Fe isomer shifts are calculated using a formalism in which the isomer shift is treated as a response of the total electronic energy with respect to the variation of the nuclear charge radius [12, 13, 14]. The isomer shift is related to the contact density as \( \delta = \alpha(\bar{\rho}_a - \bar{\rho}_s) \) where the calibration constant \( \alpha(\(^{57}\)Fe) = -0.306 ± 0.009 \) mm/s is taken from theoretical calibration of \(^{57}\)Fe isomer shift [15] and \( \bar{\rho} \) is obtained from Eq. (1).

\[
\bar{\rho}_e^{\delta/s} = \frac{5}{4\pi Z R^2} \frac{\delta E_e^{\delta/s}}{\delta R}
\]  

(1)

The derivatives \( \delta E_e(R)/\delta R \) are calculated numerically using the increment of \( 10^{-6} \) Bohr for the root mean square (r.m.s.) nuclear charge radius (\( R_{\text{Fe}} = 0.70213 \times 10^{-4} \) bohr). Throughout this work, the
3. Results and Discussion
In this section, the influence of different factors, both electronic and geometric, on the $^{57}$Fe contact densities of RbMn[Fe(CN)$_6$]·H$_2$O crystal are presented and discussed in detail. The main goal is to determine the origin of the different Fe sites observed in the HT and LT phases (see Table 1).

Table 2. $^{57}$Fe electron contact densities and isomer shift differences obtained at different theory levels.

|          | Fe$^{II}$ | Fe$^{III}$ | $\Delta \rho$ | $\Delta \delta$ |
|----------|-----------|------------|---------------|-----------------|
| HF       | 14938.17  | 14940.12   | -1.95         | 0.60            |
| MP2      | 14940.16  | 14939.82   | 0.35          | -0.11           |
| B2-PLYP  | 15140.18  | 15140.89   | -0.71         | 0.22            |
| CCSD     | 14940.38  | 14940.70   | -0.52         | 0.16            |

RbMn[Fe(CN)$_6$]·H$_2$O, iron can be present either in the Fe$^{II}$ or Fe$^{III}$ oxidation states depending on the charge transfer between iron and manganese. In this work, we have carried out calculations on both oxidation states of iron incorporated in clusters described in the previous section. In both oxidation states, iron is assumed to be present in a low-spin state: spin 0 in Fe$^{II}$ clusters and spin 1/2 in Fe$^{III}$ clusters.

Table 2 gives the $^{57}$Fe electron contact densities and isomer shifts obtained at different levels of theory (type I coordination sphere shown in Fig. 1 is assumed in these calculations). From the earlier theoretical works on K$_4$Fe(CN)$_6$ and K$_3$Fe(CN)$_6$, $\Delta \delta$ (Fe$^{II}$ - Fe$^{III}$) is expected to be positive. The Hartree-Fock, B2-PLYP and CCSD yield positive $\Delta \delta$ values while MP2 yields a negative value (see Table 2). From Table 2, it is obvious that the $\Delta \delta$ values obtained with the use of the methods including electron correlation are in much better agreement with the experimental observations than the HF value ($\Delta \delta_{\text{CCSD}} = 0.16$ mm/s and $\Delta \delta_{\text{B2-PLYP}} = 0.22$ mm/s vs. $\Delta \delta_{\text{HF}} = 0.60$ mm/s). The values obtained from the CCSD and B2-PLYP calculations are very close to one another. Therefore B2-PLYP is employed in the further calculations as a cost effective alternative to the coupled cluster method. Table 3 gives the contact densities and isomer shifts on iron in Fe$^{II}$ and Fe$^{III}$ oxidation states, for two different distribution patterns of Rb around Fe (see Fig. 1). In both types of Rb distribution patterns, change in the oxidation state of iron leads to a large variation of the isomer shift, of the order of $\Delta \delta$ (Fe$^{II}$ - Fe$^{III}$) $\sim$ 0.2 mm/s. These large variations in $\delta$ of iron sites with iron in Fe$^{II}$ and Fe$^{III}$ oxidation states agree qualitatively with the experimentally observed variations of the isomer shift in HT and LT phases. Thus, it can be concluded that iron in the HT phase is present predominantly in the form of Fe$^{III}$, whereas in the LT phase it is in the form of Fe$^{II}$.

Changing the distribution pattern of Rb$^+$ ions (see Fig. 1) leads to much smaller variations of the isomer shift for both oxidation states of iron (see Table 3). The $\Delta \delta$ values reported in the last
column of Table 3 are consistent with the isomer shift differences of $\Delta \delta_{HT} = 0.012$ mm/s and $\Delta \delta_{LT} = 0.034$ mm/s observed for different iron sites within the HT phase and the LT phase respectively (see Table 1). Thus, these differences can be interpreted as originating from differences in the Fe$^{III}$ ions distribution patterns around the Fe$^{III}$ sites in the HT phase and around the Fe$^{II}$ sites in the LT phase.

In the calculations reported above, it was assumed that the geometry of the coordination sphere of iron ions is the same in the HT and in the LT phases. In the HT phase, crystal structure of which is known experimentally, the Fe$^{III}$ ions are surrounded by six cyanido ligands at a distance $R_{Fe-C} = 1.929$ Å. Because changing the oxidation state of iron to Fe$^{II}$ should lead to contraction of the Fe-C bond length [17, 18], we have studied the effect of the bond length contraction on the contact densities and isomer shifts. In this study it was found that the bond length contraction in Fe$^{II}$ coordination sites leads to a decrease in the isomer shift difference. Therefore, the value of $\Delta \delta(Fe^{II} - Fe^{III}) = 0.22$ mm/s obtained for the type I coordination at the HT experimental geometry should be corrected to a lower value (e. g. $\Delta \delta \approx 0.09$ mm/s for Fe$^{II} - C$ bond length of 1.89 Å), thus improving the agreement with the experiment (see Table 1) [3].

4. Conclusion

The origin of the different iron sites observed experimentally in the $^{57}$Fe Mössbauer spectra of Prussian blue analogue RbMn[Fe(CN)$_6$]·H$_2$O is analyzed with the help of theoretical calculations of the isomer shift on $^{57}$Fe. In these calculations, a new formalism in which the isomer shift is treated as a response of the total electronic energy with respect to the variation of the nuclear charge radius (see Eq. (1)) is employed [12, 13, 14]. The relativistic calculations have been carried out at different levels of description of electron correlation varying from the Hartree-Fock method to the coupled cluster method (CCSD) and to the double hybrid density functional method (B2-PLYP) of Grimme[5].

The theoretically obtained $^{57}$Fe isomer shift values enable one to interpret the experimentally observed $\Delta \delta_{HT-LT}$ isomer shift differences (ca. 0.1 mm/s, see Table 1) between the HT phase and the LT phase of the RbMn[Fe(CN)$_6$]·H$_2$O compound as originating from the change of the oxidation state of iron from Fe$^{III}$ to Fe$^{II}$. Two different iron sites observed within the HT and LT phases separately, which lead to $\Delta \delta$ values of the order of 0.01-0.03 mm/s (see Table 1), can be interpreted as originating from different distribution patterns of the Rb$^+$ ions around the iron sites as shown in Fig. 1. The theoretical calculations yield the $\Delta \delta$ values of the order of 0.05 mm/s for the two types of Rb$^+$ coordination thus corroborating the analysis of the experimental measurements presented by Vertelman et al. [3].

References

[1] Sato O, Iyoda T, Fujishima A and Hashimoto K 1996 Science 272 704.
[2] Tokoro H, Ohkoshi S -L and Hashimoto K 2003 Appl. Phys. Lett. 82 1245.
[3] Vertelman E J M, Lummen T T A, Meetsma A, Bouwkamp M W, Molnar G, van Loosdrecht P H M and van Koningsbruggen P J 2008 Chem. Mater. 20 1236-38.
[4] Bartlett R J and Musial M 2007 Rev. Mod. Phys. 79 291-352 and references cited therein.
[5] Grimme S 2006 J. Chem. Phys. 124 034108.
[6] Ahlrichs R and May K Phys. Chem. Chem. Phys. 2000, 2, 943.
[7] Petersson G A and Al-Laham M A 1991 J. Chem. Phys. 94 6081-90 and references cited therein.
[8] Fuentelaña P, von Szentpaly L, Preuss H and Stoll H 1985 J. Phys. B: At. Mol. Phys. 18, 1287.
[9] a) Dyall K G 2001 J. Chem. Phys. 115 9136-43 b) Dyall K G 2002 J. Comput. Chem. 23 786-93.
[10] Dyall K G 1997 J. Chem. Phys. 106 9618-26.
[11] Filatov M and Dyall K G 2007 Theor. Chem. Acc. 117 333-38.
[12] Filatov M 2007 J. Chem. Phys. 127 084101.
[13] Kurian R and Filatov M 2008 J. Chem. Theory. Comput. 4 278-85.
[14] Kurian R and Filatov M 2009 J. Chem. Phys. 130 124121.
[15] Filatov M and Kurian R (manuscript in preparation).
[16] a) Visscher L and Dyall KG 1997 At. Data Nuc. Data Tables 67, 207. b) Visser O, Aerts PJ C, Hegarty D and Nieuwoort W C 1987 Chem. Phys. Lett. 134, 34-38.
[17] Neese F 2002 Inorg. Chem. Acta 337 181-92.
[18] Nieuwoort W C, Post D and van Duijnen P Th 1978 Phys. Rev. B 17 91-98.