REVIEW

Playing quantum hide-and-seek with the muon: localizing muon stopping sites

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Abstract

The most fundamental limitations of a muon-spin relaxation experiment can be the lack of knowledge of the implantation site of the muon and the uncertainty about the muon’s perturbation of its host. Here we review some of the work done on the ‘muon site problem’ in the solid state and highlight some recent applications of electronic structure calculations that have successfully characterized the quantum states of muons in a number of insulating compounds containing fluorine, in a number of pnictide superconductors, and in ZnO.

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

1. Introduction

A muon-spin relaxation (μSR) experiment involves implanting spin-polarized positive muons in a sample in order to probe the local static and dynamic magnetic properties (a more detailed introduction can be found in [1] and in the lead article of this series [2]). μSR is an extremely sensitive probe of magnetism (for an illustration see e.g. [3]) but it has two significant limitations. The first concerns the lack of knowledge of the site of implantation of the muon, which hinders the measurement of magnetic moments or the comparison of different candidate magnetic structures using μSR. Second, the unknown extent of the perturbation due to the muon of the local crystal and electronic structure of the host has been the cause for increased concern since μSR is frequently employed in the study of systems that lie on the verge of ordering [3–6] or where doping is a critical parameter [7–9].

From the very beginning of μSR, significant effort has been devoted to the determination of muon sites. In some materials a determination of interstitial muon sites was indeed possible thanks to accurate experimental studies of the muon frequency shift in an applied magnetic field [10], level crossing resonances [11, 12], by inspecting relaxation rates as a function of applied field [13, 14] or through the observation of quantum entanglement between the muon...
spin and a small number of surrounding nuclei (discussed in more detail below) [15, 16]. Nonetheless the number of examples where the muon site can be determined by experimental means alone is limited and even in those cases the experimental information about the muon site and the perturbation caused by the muon is usually incomplete. An improved understanding of the muon state in solids would not only benefit a more complete understanding of the nature of the muon response in a wide range of compounds but also it could enable a determination of magnetic moments and perhaps even allow to differentiate between different models of magnetic structures. This information would be particularly valuable in many topical compounds where the observation of magnetic neutron scattering is challenging, such as compounds containing nuclei that strongly absorb neutrons (e.g. iridates) or compounds with particularly small magnetic moments (e.g. frustrated and low-dimensional systems, where the moments are strongly renormalized by quantum fluctuations).

In this review we present three case studies that characterize the muon states in solids using \textit{ab initio} electronic structure theory. Previous work in this area has focused on the paramagnetic states formed by muons and protons in semiconductors, for a review see [17]. Diamagnetic muon states (where the contact hyperfine coupling is negligible) have received very little attention in spite of their greater utility in the study of magnetic materials. \textit{Ab initio} methods have also been applied in the study of muoniated molecular radicals, which is the subject of another article in this series [18]. Here we summarize a number of recent applications of density-functional theory (DFT) that focus on the diamagnetic muon states in a number of solids. The paramagnetic muonium states in some of the fluorides discussed here have also been studied in some detail (including a calculation of the contact hyperfine couplings and the quantum corrections to it). The interested reader is referred to [19] for the details.

2. Quantum states of muons in insulating fluorides

In host compounds containing fluorine, diamagnetic muons can couple strongly to the fluoride ions, often forming linear F–μ–F complexes [15], although bent F–μ–F and F–μ geometries have been shown to exist as well [16]. The magnetic dipolar coupling between muon and fluorine nuclear spins $I$ (both $I = 1/2$) is described by the Hamiltonian

$$\mathcal{H} = \sum_{i>j} \frac{\mu_0 \gamma_f}{4\pi r^3} \left[ I_i \cdot \mathbf{r} \cdot \hat{r} (I_j \cdot \hat{r}) \right],$$

where $\hat{r}$ is the normalized vector connecting spins $i$ and $j$, $\gamma_f$ is the gyromagnetic ratio of spin $i$, $r$ is the distance between spins $i$ and $j$ and all other symbols take their usual meaning. This interaction gives rise to a characteristic muon precession signal (which can be easily determined by diagonalizing the Hamiltonian) that is sensitive to the geometry of the muon–fluorine state, allowing an accurate experimental determination of the muon’s local site geometry [15, 16]. Two recent studies [19, 20] have investigated the quantum states of muons in the non-magnetic ionic insulators LiF and NaF (rock-salt structure), CaF$_2$ and BaF$_2$ (fluorite structure), CoF$_2$ (orthorhombic) and for the antiferromagnetic insulator CoF$_2$ (rutile-type structure).

![Figure 1](image.png)

Figure 1. Calculated equilibrium geometries of dia- and paramagnetic muonium states in LiF/NaF (Li/Na blue and F green), CaF$_2$/BaF$_2$ (Ca/Ba red) and CoF$_2$ (Co magenta). Translucent spheres represent the equilibrium ionic positions before the muon (brown) is introduced into the crystal. Black lines are a guide to the eye. The $c$-axis is vertical. Reproduced with permission from [19]. Copyright 2013 by American Physical Society.

Both studies used the plane-wave pseudopotential method as implemented in the Quantum ESPRESSO package [23]. The calculations employ periodic boundary conditions and so to reduce the error due to defects in neighbouring unit cells, a supercell approach was used where each supercell contained $2 \times 2 \times 2$ conventional unit cells (except for YF$_3$ where the conventional orthorhombic unit cell was used). In these studies, the charge state of the muon was determined by the charge of the supercell (+1 for diamagnetic and neutral for paramagnetic states). Two alternative approaches were used for determining the relaxed geometries: the first placed a muon in several randomly chosen low-symmetry sites and all ions were allowed to relax until the forces on all ions and the energy change between iterations had fallen below a convergence threshold [19]; the second calculated the electrostatic potential of the unperturbed solid first, placed muons in the local minima of the electrostatic potential, and then the structure was allowed to relax [20].

Figures 1 and 2 show the calculated equilibrium geometries of the muon states in the compounds considered. In all cases, an F–μ–F state is the lowest energy state. The F–μ bond length of about 1.18 Å (calculated value for the
Based on figures in bond between the muon and the nearest neighbour (nn) F atoms. Previously suggested linear F–µ instead, a bent F–µ the ground state diamagnetic site of the muon in YF₃ sites for the muon. On kinetic grounds they predict that their DFT calculations, the authors found several candidate µ was attributed to the formation of a linear F–µ configuration with the results obtained using the procedure outlined in the text, which predicts a slightly distorted bond between the muon and the nearest neighbour (nn) F atoms. Based on figures in [20]. The data were visualized with VESTA [22].

(F–µ–F)⁻ (molecule in vacuum) corresponds to approximately one ionic radius of the fluoride ion. The (F–µ–F)⁻ molecule can therefore be pictured as two hard spheres of fluoride ions that are just touching with the muon in between. In LiF, NaF, CaF₂ and BaF₂ the calculations correctly predict the experimentally known geometries [15] with great accuracy: bond lengths are within ~3% of the experimental values. Even though the muon site in CoF₂ agrees with the site known from a detailed experimental study [10], the predicted F–µ–F state had not been observed experimentally. Following their ab initio work, the authors experimentally searched for signatures of an F–µ–F state in CoF₂ and found unambiguous evidence for a linear F–µ–F state of a geometry that is in excellent agreement with their DFT prediction [19]. In YF₃ an F–µ–F signal had been previously observed [21] and was attributed to the formation of a linear F–µ–F state. In their DFT calculations, the authors found several candidate sites for the muon. On kinetic grounds they predict that the ground state diamagnetic site of the muon in YF₃ is, instead, a bent F–µ–F state with a bond angle of about 144°. They revisit the previous experimental data and show that the crystallographic distortions are significant at short range, with nn fluoride distortions of approximately 0.25 Å (LiF), 0.5 Å (NaF and CoF₂), 0.2 Å (CaF₂) and 0.4 Å (BaF₂) and nn cation distortions of 0.4 Å (LiF, NaF, and BaF₂), 0.35 Å (CaF₂) and 0.2 Å (CoF₂). While it was known that the perturbation of the fluoride ions must be significant based on the experimentally measured F–µ bond lengths of the F–µ–F states found in many fluorides, these calculations allow the cation distortions to be quantified as well. Since localized magnetic moments would be located on the cation, the cation displacements are particularly pertinent to understanding the effect of the muon’s perturbation on experimentally measured µ⁺SR spectra. The results above demonstrate that the perturbation of the nn cations can even exceed those of the fluoride ions bound in the F–µ–F state. In antiferromagnetic CoF₂ they calculate that the crystallographic distortion of the Co cations will lead to a reduction of the observed muon precession frequency by just over 20%, in good agreement with an estimate based on experimental data [10] of 16%. This correction should be taken into account if magnetic moments were to be measured accurately in an ionic insulator. Since at short distances the distortions are mainly caused by the electrostatic interaction of the unscreened muon with its surroundings, the authors expect similar distortions in any ionic insulator. While the muon is more screened in more covalent compounds, it was recently shown for the organic radical magnets F₄BImNN and BImNN [24] that the relative ‘softness’ of covalent bonds can lead to similar crystallographic distortions.

The quantum nature of nuclei is generally ignored in DFT calculations since nuclear masses are typically so large that quantum effects (caused e.g. by the spread of the nuclear wavefunction) lie below the current level of accuracy of the technique. However, at approximately 1/9 the mass of a proton the muon is an exceptionally light impurity. Quantum effects can therefore be expected to play a more significant role in the localization of a muon than for the majority of conceivable point defects. The quantum properties of the muon in the F–µ–F state were studied using density-functional perturbation theory by calculating the vibrational properties of the F–µ–F molecule. The zero-point energy (ZPE) was then estimated from the calculated vibrational frequencies in the harmonic approximation. This approximation neglects the finite spread of the muon wavefunction and anharmonic terms in the potential. However, it takes account of the coupled muon–ion zero-point motion and is most appropriate for a molecular defect such as the F–µ–F system. In fact it was found that in most of these systems the vibrational modes of the F–µ–F centre decouple from the rest of the crystal and so the F–µ–F system can be viewed as a molecule-in-a-crystal defect. The muon–fluoride (or hydrogen-fluoride (HF)) bond is the strongest known hydrogen bond in nature [25] with a bond enthalpy of 565 kJ mol⁻¹ (for HF) compared with 463 kJ mol⁻¹ for the hydroxyl bond in OH [26]. Combined with the small muon mass this leads to the F–µ–F centre possessing an exceptionally large ZPE: larger than that of any natural triatomic molecule [19]. This demonstrates the importance of quantum effects on muon localization.

There will also be a quantum correction to the dipolar coupling, which has not been included at this level of
approximation. While a direct spin relaxation due to the quantum zero-point motion is motionally narrowed out from the spectra, such that only the average dipolar coupling is observed, there will be a quantum correction to this average due the finite muon nuclear wavefunction spread. There will also be a small shift in inter-atomic distances due to the muon (or proton) zero-point motion. This effect was estimated to be on the order of 0.03 Å for the HF bond length in the (FHF)$^-$ ion [27] (see below). The already good agreement of the calculated dipolar coupling with the experimental value demonstrates that this quantum correction will be small in this case. However, in a situation where spin density were to leak onto the muon as a result of its zero-point motion, there may be a sizeable correction to the magnetic coupling of the muon.

The quantum treatment of the muon discussed above is applied after solving the problem for a classical (i.e. infinitely heavy muon). It is worth noting that some attempts have been made to take account of the quantum nature of the muon (or proton) from the outset. One approach, called nuclear-electronic orbital (NEO) method, based on second-order Møller–Plesset perturbation theory (MP2) [27], was applied to the (FHF)$^-$ ion with good results. The HF bond length in the (FHF)$^-$ ion was estimated to increase by approximately 0.03 Å due to the proton zero-point motion and wavefunction spread. Note however that (a) MP2 is a post-Hartree–Fock method not uncommonly applied in computational chemistry codes for small systems but cannot easily be applied to calculations in solids and (b) the quantum effects found for the (FHF)$^-$ ion are on the same scale as numerical differences between different functionals and the level of improvement expected from a NEO treatment of nuclear quantum effects is therefore, at least in the (FHF)$^-$ system, below the level of accuracy of DFT and is unlikely to be universal (for the (FHF)$^-$ system the bond length agrees less well with the experimental value when quantum effects are taken into account using the NEO method). A similar method to treat light nuclei such as protons and muons on the same footing as electrons was applied in Hartree–Fock calculations of vacancy-containing defect complexes in diamond with some success [28]. A full treatment of nuclear quantum effects within state-of-the-art DFT, however, is likely to be only possible within the path-integral molecular dynamics (PIMD) formalism [29]. Some attempts at using PIMD to study muonium in Si [30–32] and diamond [33] have been made.

There has been considerable interest recently in identifying muon sites by locating the minima of the electrostatic potential of the unperturbed host [34–38]. Möller et al [19] and Bernardini et al [20] have therefore compared the muons sites in this series with the location of the minima of the electrostatic potential of the unperturbed solid, and have found that these do not generally coincide (see figure 3). In the diamagnetic case this is primarily due to the formation of the molecular F–µ–F state. All of the compounds studied here are very ionic in character and the µ$^+$-lattice interaction is therefore expected to be stronger than in more covalent insulators or metals, where the µ$^+$ charge would at least be partially screened. However, if the muon charge were completely screened there would be no reason why a muon should localize in an electrostatic minimum. We expect the combination of this screening (where operative), the muon–lattice interaction, and the muon’s exceptionally large ZPE to frequently lead to muon localization away from the minima of the electrostatic potential of the unperturbed host. We therefore believe that muon sites cannot be determined reliably on the basis of the electrostatic potential alone.

### 3. Muon sites in metallic systems: pnictide superconductors

In this section we discuss the application of DFT to the determination of muon sites in pnictide superconductors. The analysis of the magnetic ground state properties by means of $\mu$SR in the pnictides triggered the interest to calculate the muon stopping sites [35, 37, 39]. It is well known that the magnetic and structural properties of pnictides are not accurately described in DFT because of electronic correlations. Nonetheless we have demonstrated that it is possible to determine muon sites that are consistent with the experimental $\mu$SR data.

Muon sites were identified by calculating the electrostatic potential of the unperturbed host. In some cases the muon...
The ground state energy $E_0$ was found by solving the Schrödinger equation for the muon in the electrostatic potential of the host (either for the unperturbed host or with relaxed ionic positions due to the presence of the muon). This ‘rigid-lattice’ approximation takes full account of anharmonic terms in the potential, but it neglects the effect of the muon on the surrounding charge density and the coupled muon–ion zero-point motion. Its greatest advantage is the much reduced complexity of the calculation compared with calculating the vibrational modes, as described above, which is approximately a factor $3N$ more computationally expensive ($N$ being the number of atoms in the supercell). It is also possible to use the total energy from a series of self-consistent calculations including the muon with different muon positions as potential for which the Schrödinger equation is solved. This would take account of the muon’s effect on the surrounding charge density, but would be costly to do for a full three-dimensional grid. In either case this approximation is most appropriate for an atomic defect such as muonium [19] or a diamagnetic muon in a screened environment such as a metal [37–39], as is the case here.

A number of recent successes [37–39] with pnictides demonstrate that the prediction based on the solution of the muon Schrödinger equation for the unperturbed lattice can be as accurate as required to understand and extract quantitatively consistent results from $\mu^+\text{SR}$ spectra. A few representative cases are reported in table 1. In figure 4 we compare the muon position in LaCoPO [38] estimated by considering the minimum of the electrostatic potential with the one obtained through a full ionic relaxation (including the muon) the procedure outlined above: the displacement from the potential minimum is approximately 0.25 Å. This displacement can have a significant effect on the calculated dipolar field at the $\mu^+$ site.

### 4. Molecular-orbital cluster and potential methods to determine the muon site and hyperfine interactions in La$_2$CuO$_4$ and ZnO

Even in La$_2$CuO$_4$, the parent compound of the family with the simplest crystal structure of high-T$_c$ compounds, the exact muon stopping site is uncertain. Based on dipole-field calculations, Hitti et al [42] have estimated the muon stopping site to be near the apical oxygen of the CuO$_2$ octahedra in La$_2$CuO$_4$, which was supported by later calculations of the electrostatic potential [43]. A different study [44] suggested the muon position to be 1 Å away from the in-plane oxygen through measurements of the nuclear dipole field distribution at the muon site, aided by a calculation of the electrostatic potential. Such a stable binding state between oxygen and muon has also been suggested by other ab initio calculations [45]. More recent work has lead to even more suggestions about the muon stopping site in La$_2$CuO$_4$ [46, 47].

In order to shed more light on this problem, Watanabe et al [48] propose to develop a strategy for identifying muon stopping sites by studying the muon sites in ZnO, where some information about the muon sites is available experimentally. ZnO is a wide-gap semiconductor that has been extensively studied due to its technological significance [49]. ZnO tends to exhibit n-type conductivity, although the source of this conductivity remains controversial. Two $\mu^+\text{SR}$ studies confirmed the existence of muonium centres in ZnO; one experiment observed a single muonium centre [50] with contact and dipolar hyperfine couplings of $A = 500 \pm 20$ kHz and $D = 260 \pm 20$ kHz, respectively, while the other study detected two signals corresponding to two distinct muonium

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**Figure 4.** Comparison of the localization volume based on the electrostatic potential of the unperturbed host (shaded yellow) and the relaxed muon site obtained through a structural relaxation (labelled $\mu^+$) in LaCoPO. Further details can be found in [38].

**Table 1.** Comparison of the experimental ($B_{\text{exp}}^\text{dip}$) and the calculated ($B_{\text{calc}}^\text{dip}$) dipolar field at the $\mu^+$ site for two parent compounds of the pnictide high-temperature superconductors. The minima of the electrostatic potential are used to evaluate $B_{\text{calc}}^\text{dip}$ on the basis of the neutron scattering results for the Fe magnetic moments reported in [40, 41] (only significant figures are reported). $\mu^+\text{SR}$ results are taken from [37, 39] which contains further details of the calculation.

| Compound           | $B_{\text{exp}}^\text{dip}$ (G) | $B_{\text{calc}}^\text{dip}$ (G) |
|--------------------|-------------------------------|----------------------------------|
| FeTe              | 2230                          | 2000(100)                        |
| LaFeAsO site 1    | 1530                          | 1650(50)                         |
| LaFeAsO site 2    | 270                           | 180(10)                          |
centres [51]. These two centres were proposed to be the so-called anti-bonding centre (AB) with $A = 491(5)$ kHz and $D = 265(9)$ kHz and the bond centre (BC) site with $A = 293(7)$ kHz and $D = 286(13)$ kHz (see figure 5). While strictly still hydrogenic, muonium in ZnO is not at all hydrogen-like. Instead, the exceptionally small hyperfine couplings indicate the highly dilated nature of the Mu electron wavefunction. This implies that screening is negligible and that the muonium site must therefore be identical to the diamagnetic site. In other words, muonium (hydrogen) acts as a shallow donor.

For ionic compounds, the customary practice to treat the boundary conditions is to embed the cluster with a finite number of point charges that would reproduce the correct Madelung potentials [52]. ZnO has both ionic and covalent character in its bonding. The usage of hydrogen to terminate the dangling bonds could therefore have significant effects on the electronic structure, especially for the BC site. To examine the effects of hydrogen terminating dangling bonds in the molecular-orbital (MO) cluster method, Watanabe et al have performed MO cluster calculations for muonium at the BC site with and without the hydrogen terminators. The clusters contained eight Zn and O atoms and one hydrogen to represent the muonium. For muonium at the BC site, Watanabe et al found that the lattice relaxation effect is about 40% which is consistent with a previous ab initio study [49].

Watanabe et al have also employed both Hartree–Fock and DFT calculations (the PBE and B3LYP functionals were used with similar results) to calculate the hyperfine coupling constants for muonium at the BC site using the Gaussian 03 software. They found that by not using hydrogen as terminators, the hyperfine coupling constants were reduced significantly. Comparing to the results of Li et al [53], the isotropic Fermi contact coupling constant $A$ is reduced by a factor of 35 while for the dipolar component $D$ the reduction is by a factor of 56. For $A$, the sign remained negative both for clusters with and without hydrogen terminators. The hyperfine coupling constants calculated using DFT were smaller than those obtained previously [53] and the sign of $A$ was positive in Watanabe et al’s work, in agreement with the experimental data and improving on previous work [53]. Further work will investigate whether embedding the cluster in an assembly of point charges would further bring the values of the hyperfine coupling constants closer to the experimental ones.

5. Conclusions

We have discussed the motivation for investigating the location of muon sites and the extent of the perturbation caused by the muon. We have presented recent successes in the study of muon states in wide-gap insulating fluorides, where the local muon site can be determined experimentally with high accuracy, in pnictide superconductors, and in ZnO. These results demonstrate that DFT is a powerful tool to characterize muon states in a wide range of solids. With the continuing improvements of electronic-structure methods and the growing performance of the computational resources available, muon states can be explored more accurately and in greater detail than ever before, even in challenging materials. We believe that this will become a routine part of many muon experiments and will boost the range of physical properties that can be explored with $\mu^+$SR. This work is supported by EPSRC (UK), the European 7th framework programme contract 226507 (NMI3), RIKEN (Japan) and Universiti Sains Malaysia.

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