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Understanding the μ SR spectra of MnSi without magnetic polarons

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Transverse-field muon-spin rotation (μ SR) experiments were performed on a single crystal sample of the noncentrosymmetric system MnSi. The observed angular dependence of the muon precession frequencies matches perfectly the one of the Mn-dipolar fields acting on the muons stopping at a $4a$ position of the crystallographic structure. The data provide a precise determination of the magnetic dipolar tensor. In addition, we have calculated the shape of the field distribution expected below the magnetic transition temperature T_C at the $4a$ muon site when no external magnetic field is applied. We show that this field distribution is consistent with the one reported by zero-field μ SR studies. Finally, we present *ab initio* calculations based on the density-functional theory which confirm the position of the muon stopping site inferred from transverse field μ SR. In view of the presented evidence we conclude that the μ SR response of MnSi can be perfectly and fully understood without invoking a hypothetical magnetic polaron state.

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I. INTRODUCTION

In recent years a growing interest has been focused on crystalline systems lacking inversion symmetry (see, for example, Ref. [1]). A direct consequence of noncentrosymmetry is the occurrence of a spin-orbit Dzyaloshinsky-Moriya (DM) interaction [2,3]. The DM interaction is thought to play an important role in multiferroic systems and may be at play in the occurrence of nontrivial magnetic structures. On the other hand, noncentrosymmetric systems presenting superconductivity have attracted special attention as the absence of parity symmetry considerably restricts the possible superconducting states (i.e., a pure triplet pairing is in principle not allowed without inversion symmetry) [4].

MnSi is probably the noncentrosymmetric system which concentrates most of the research focus. It was the first material found to exhibit a homochiral spin spiral structure below $T_C \simeq 29.5$ K [5–7]. This spin spiral adopts a very long wavelength of about 18 nm. In addition to an external pressure destroying the magnetic state ($p_c \simeq 1.5$ GPa [8]), a modest magnetic field of 6 kG is found to overcome the DM interaction leading to a spin-aligned state [9]. Moreover a skyrmion phase was recently identified near T_C in the magnetic phase diagram [10]. Contrary to what was observed in a number of strongly correlated electron systems (see, for example, Ref. [11]), MnSi does not display superconductivity at p_c . It was invoked [12] that the absence of a superconducting state for MnSi under pressure has to be traced back to the absence of a parity operator for the wave functions.

In view of all these properties, MnSi has been investigated several times by μ SR in zero or externally applied pressures

(see, for example, Refs. [13–17]). Two main controversies occurred from these studies. The first one regards the behavior of the magnetic state near the critical pressure p_c . First measurements performed by Uemura *et al.* [16] seemed to show a phase separation between magnetic and paramagnetic regions whereas measurements performed by Andreica *et al.* [17], extending at lower temperatures, showed a complete absence of phase separation. The second controversy regards the muon stopping site(s) and the occurrence or not of magnetic polaron. The large majority of the studies associates the two spontaneous frequencies seen below T_C to two magnetically inequivalent sites. The same is deduced from transverse-field (TF) measurements performed at low fields [18], even though the conclusions obtained from this latter study are blurred by the assumption of a vanishing dipolar tensor. On the other side, Storchak *et al.* [19] relate the observation of two frequencies in TF data to a coupled μ^+e^- system (i.e., Mu state). In this scenario, the spin of the Mu electron does not exhibit rapid fluctuations through spin exchange with the itinerant electrons, but is kept “fixed” by the local ferromagnetic ordering mediated by itself (the electron and the four Mn neighbors are thought to be coupled and to behave as a single entity with large spin: i.e., a “spin polaron”). In this scenario, the two frequencies correspond to two muon spin-flip transitions between states with fixed electron spin orientation. The picture of such a localized electron state appears controversial given the metallic nature of MnSi.

The aim of the present paper is to lift the controversy regarding the occurrence or not of magnetic polaron and provide information on the muon stopping site. We retain this point essential to solve, as this will show whether μ SR can be utilized to specifically extract quantitative information on this heavily studied strongly correlated system. In this study, we will show that the occurrence of different frequencies

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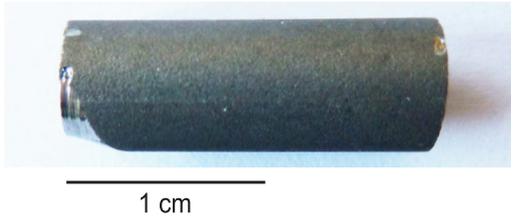


FIG. 1. (Color online) Sample used for the TF- μ SR measurements. The sample was rotated along the cylinder axis.

in the TF- μ SR spectra can be traced back to the different responses of crystallographically equivalent muon-stopping sites which become magnetically inequivalent in the presence of an external magnetic field and that one does not need to invoke a magnetic polaron to explain the TF- μ SR data. We will also show that the occurrence of two spontaneous muon frequencies in the zero-field μ SR data recorded below T_C can be explained within this frame. Finally, we also present density-functional theory calculations which perfectly support the experimental evidences.

II. EXPERIMENTAL DETAILS

We used the same MnSi sample as in the previous μ SR under pressure experiments [17]. It consists of a single crystal with the form of a cylinder of approximately 7 mm diameter and 19 mm length (see Fig. 1). The single crystal was grown by the Czochralsky pulling technique from a stoichiometric melt of high-purity elements (>99.995%) using radio-frequency heating and a cold copper crucible. No deviation from the known crystal structure [space group $P2_13$, No. 198; Mn-ion at the position (0.138,0.138,0.138), and Si-ion at the position (0.845,0.845,0.845); lattice constant 4.558 Å] nor any presence of foreign phase was detected by scanning electron microscope microanalysis, backscattered electron imaging, and the x-ray Debye Scherrer pattern.

The study of the angular dependence of the muon Knight shift in MnSi was performed using the GPS instrument located at the π M3 beamline of the HIPA Complex at the Paul Scherrer Institut (PSI, Villigen, Switzerland). The measurements were performed with the transverse-field geometry with a magnetic field of 5200 G. The sample was rotated around the cylinder axis and the external field \mathbf{B}_{ext} was applied perpendicular to the cylinder. Note that the sample was not grown along a specified crystallographic orientation, i.e., the cylinder axis does not correspond to a principal axis. The measurements presented here were performed using a dynamical He-flow cryostat (i.e., sample in He-flow) at a temperature of 50 K. The temperature stability of the sample was better than 0.1 K for all the measurements.

III. RESULTS AND DISCUSSION

A. Transverse-field data and muon site determination

For all rotation angles ϕ' , the μ SR signals are best modeled by the presence of four frequencies with different angular

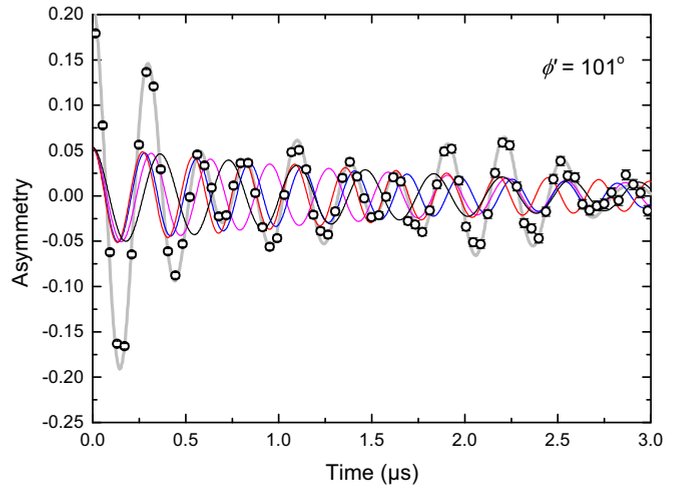


FIG. 2. (Color online) Example of one μ SR time spectrum plotted in a rotating reference frame (RRF) with frequency 65 MHz. The gray line represents the fit to the data (open points) using Eq. (1) and the four individual signals are also displayed (color code is the same as in Fig. 4). Note that the actual fits were performed without RRF and without time binning (i.e., original bin of 0.9766 ns). Notice that the four signals have very similar depolarization rates.

dependences. Fits with all parameters free point to similar amplitudes for the four components. Hence, in a second series of fits the amplitudes were forced to be the same. That is,

$$\mathcal{A}_{\text{tot}}G(t) = \sum_{i=1}^4 \mathcal{A} \exp(-\lambda_i t) \cos(v_i 2\pi t + \psi). \quad (1)$$

The frequencies v_i reflect the local magnetic field values sensed by the muon at the stopping site(s), i.e., $v_i = \gamma_\mu |\mathbf{B}_{\text{loc,TF},i}| / (2\pi)$, where γ_μ is the gyromagnetic ratio of the muon. The phase parameter ψ is common to all signals as it is defined by the direction of the muon spin with respect to the detector system at the muon implantation time. During the fits, no restrictions were applied concerning the frequencies and the results were directly obtained through the software MUSRFIT [20]. Note that all the fits were performed in the time space between 0 and 8 μ s. As illustrations to the data, we present in Fig. 2 a μ SR time spectrum, plotted in a rotating reference frame, and in Fig. 3 selected fast Fourier transform (FFT) of spectra recorded at different angles. The full angular dependence of the frequencies is shown in Fig. 4.

The occurrence of four frequencies in the μ SR signal clearly points to a muon site located at the $4a$ Wyckoff position of the cubic structure. In this view, the main contribution of the observed angular dependence arises from the dipolar contribution. As shown in Table I, which provides information on this type of site and the respective dipolar tensor forms, taking into account the symmetry of the dipolar tensors, the four crystallographically equivalent $4a$ sites in the unit cell are either (i) magnetically equivalent when \mathbf{B}_{ext} is applied along a principal direction; (ii) grouped in two pairs of magnetically inequivalent sites for \mathbf{B}_{ext} applied in a principal plane; (iii) usually all magnetically inequivalent for \mathbf{B}_{ext} applied along an arbitrary direction.

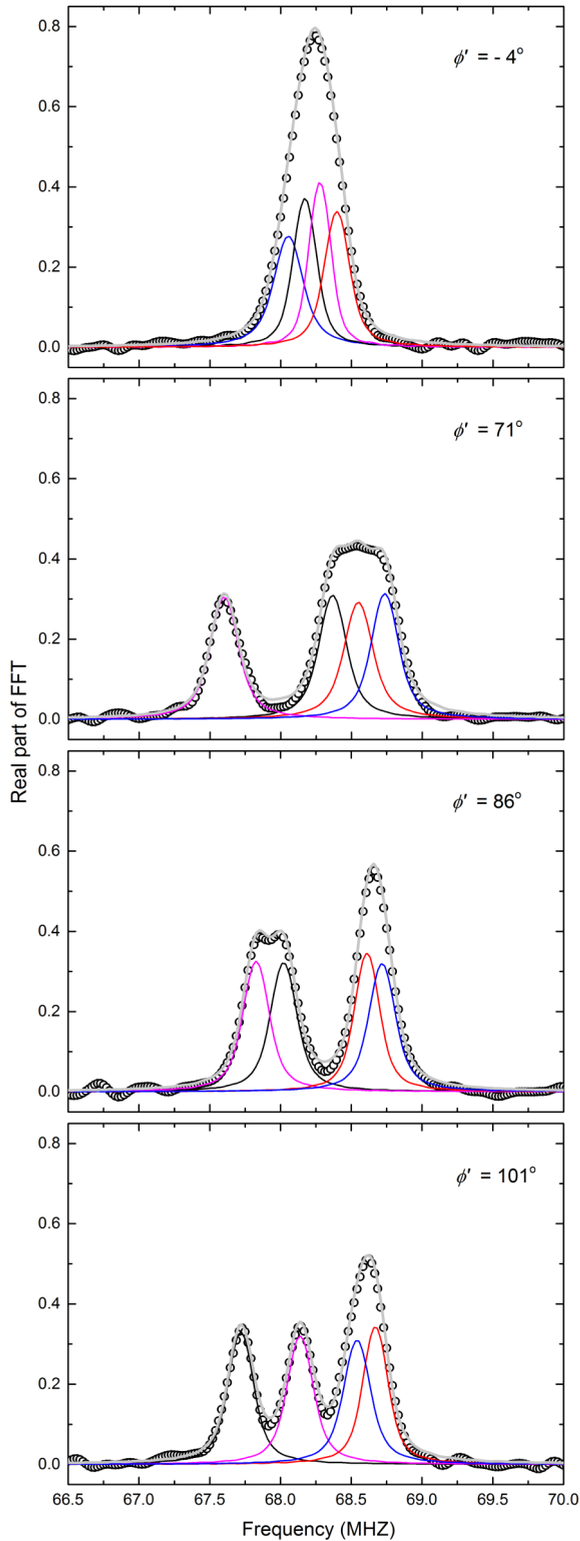


FIG. 3. (Color online) Fourier transform of TF spectra taken at different orientations. The color code of the different signals is the same as in Figs. 2 and 4.

Generally, the local field at the muon site can be written as usual as (see, for example, Refs. [21,22])

$$\mathbf{B}_{\text{loc,TF}}(\phi') = \mathbf{B}_{\text{ext}} + \mathbf{B}_{\text{cont}} + \mathbf{B}_{\text{dip}}(\phi') + \mathbf{B}_{\text{Lor}} + \mathbf{B}_{\text{dem}}(\phi'), \quad (2)$$

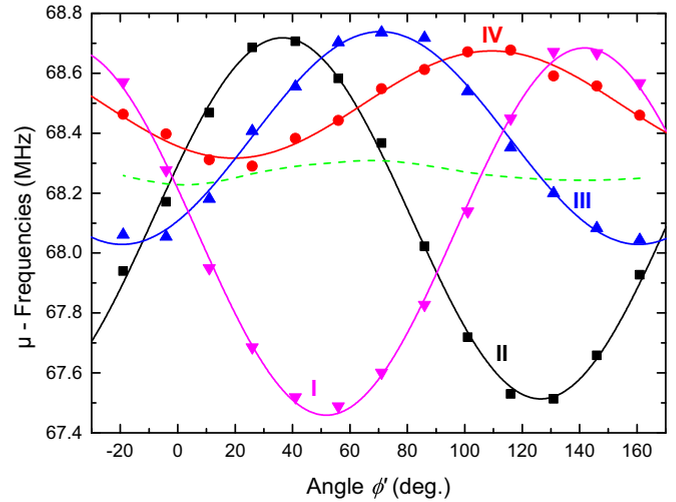


FIG. 4. (Color online) Angular dependence of the fitted μ SR frequencies. The lines are guides to the eye. The green dashed line represents the average frequency $\bar{\nu}(\phi')$ (see text).

where the values of \mathbf{B}_{ext} , of the contact term \mathbf{B}_{cont} , and of the Lorentz field \mathbf{B}_{Lor} are independent of the rotation angle ϕ' (bear in mind that for a cubic system, the representation of the magnetic susceptibility tensor in the crystal reference frame is given by $\chi = \chi \mathbf{E}$, where $\mathbf{E} = [E_{ij}] = [\delta_{ij}]$). We mention at this point that, as said, we expect that the essential part of the observed angular dependence should arise from the dipolar contribution. Nevertheless, we cannot exclude a small angular dependence of the demagnetization field, as the sample had not a perfect cylindrical shape (see Fig. 1). A way out is to observe that (i) for a given field direction the demagnetization field is identical for all signals; (ii) due to the symmetry of the dipolar tensors for the $4a$ sites, the sum of the dipolar contribution of the four signals should always cancel (see Table I). Hence the true angular dependence of the measured dipolar contribution for each signal can be obtained by

$$B_{\text{dip},i}(\phi') = \frac{(\nu_i(\phi') - \bar{\nu}(\phi')) 2\pi}{\gamma_\mu}, \quad (3)$$

where $\bar{\nu}(\phi')$ is the average frequency at each angle. We observe that the amplitude of the angular dependence of $\bar{\nu}(\phi')$ is very small, i.e., it represents a field $\bar{\nu}(\phi') 2\pi / \gamma_\mu$ of the order of 1.8 G, which would correspond to a variation of less than 3% of the demagnetization factor during the rotation (see Fig. 4). The average frequency $\bar{\nu}(\phi')$ can be interpreted as the sum of all other contributions, that is,

$$\bar{\nu}(\phi') = \frac{\gamma_\mu}{2\pi} |\mathbf{B}_{\text{ext}} + \mathbf{B}_{\text{cont}} + \mathbf{B}_{\text{Lor}} + \mathbf{B}_{\text{dem}}(\phi')|. \quad (4)$$

For the four signals, the dipolar contributions obtained with the slight correction described in Eq. (3) are shown on Fig. 5.

In the following, we will carefully analyze the angular dependence of these dipolar contributions to determine the muon-stopping site. The dipolar field contribution to the value of the local field sensed by the muon can be written as

$$\mathbf{B}_{\text{dip}} = \mathbf{A}_{\text{dip}} \chi \mathbf{B}_{\text{ext}}, \quad (5)$$

TABLE I. Coordinates and form of the representation of the dipolar tensor \mathbf{A}_{dip} in the crystal reference frame for the crystallographic Wyckoff position $4a$ (four crystallographically equivalent sites designed here by I to IV). We shall call $[A_{ij}^c]$ this representation. Note that the value $|a_{\text{dip}}|$ is the same for all the tensors. Bear also in mind that the structure can also accommodate Wyckoff positions of type $12b$ (all other sites not equivalent to a $4a$ position) but in this case one would expect to observe 12 frequencies in the μSR signal for an arbitrary direction of the applied magnetic field.

	Site 4a-I	Site 4a-II	Site 4a-III	Site 4a-IV
Coordinates	(x, x, x)	$(\frac{1}{2} - x, \bar{x}, \frac{1}{2} + x)$	$(\frac{1}{2} + x, \frac{1}{2} - x, \bar{x})$	$(\bar{x}, \frac{1}{2} + x, \frac{1}{2} - x)$
$[A_{ij}^c]$	$\begin{pmatrix} 0 & a_{\text{dip}} & a_{\text{dip}} \\ a_{\text{dip}} & 0 & a_{\text{dip}} \\ a_{\text{dip}} & a_{\text{dip}} & 0 \end{pmatrix}$	$\begin{pmatrix} 0 & a_{\text{dip}} & -a_{\text{dip}} \\ a_{\text{dip}} & 0 & -a_{\text{dip}} \\ -a_{\text{dip}} & -a_{\text{dip}} & 0 \end{pmatrix}$	$\begin{pmatrix} 0 & -a_{\text{dip}} & -a_{\text{dip}} \\ -a_{\text{dip}} & 0 & a_{\text{dip}} \\ -a_{\text{dip}} & a_{\text{dip}} & 0 \end{pmatrix}$	$\begin{pmatrix} 0 & -a_{\text{dip}} & a_{\text{dip}} \\ -a_{\text{dip}} & 0 & -a_{\text{dip}} \\ a_{\text{dip}} & -a_{\text{dip}} & 0 \end{pmatrix}$

where \mathbf{A}_{dip} is the dipolar tensor and $\chi \mathbf{B}_{\text{ext}}$ represents the local Mn moment induced by the external field. As in the present experiment the field was not rotated in a principal crystallographic plan, we will define $[A_{i'j'}^l]$ as the representation of the dipolar tensor in the reference frame given by the rotation axis and rotation plane of the applied field (i.e., the laboratory frame; see Fig. 6). Expressing the rotation axis with the Euler angles θ and ϕ , we have the relation,

$$[A_{i'j'}^l] = R_{[010],\theta} R_{[001],\phi} [A_{ij}^c] R_{[001],-\phi} R_{[010],-\theta}, \quad (6)$$

where the rotation matrices $R_{\alpha,\beta}$ represent a rotation of angle β around the direction α and $[A_{ij}^c]$ is the representation of the dipolar tensor in the crystal reference frame (see Table I). In the reference frame of the rotation axis and plane, we can express the external field as $\mathbf{B}_{\text{ext}} = B_{\text{ext}}(\sin \theta' \cos \phi', \sin \theta' \sin \phi', \cos \theta')$, where θ' and ϕ' are now the polar and azimuth angles of the field in the new reference frame (see Fig. 6). Generally, the angular dependence of the

component of \mathbf{B}_{dip} along the direction of the external field can be expressed as [23]

$$\begin{aligned} B_{\text{dip},\parallel} = & \frac{1}{3}(A_{x'x'}^l \chi_{x'}^l + A_{y'y'}^l \chi_{y'}^l + A_{z'z'}^l \chi_{z'}^l) B_{\text{ext}} \\ & + \frac{2}{3}[A_{z'z'}^l \chi_{z'}^l - \frac{1}{2}(A_{x'x'}^l \chi_{x'}^l + A_{y'y'}^l \chi_{y'}^l)] P_2^0(\cos \theta') B_{\text{ext}} \\ & - \frac{1}{3}A_{x'z'}^l (\chi_{x'}^l + \chi_{z'}^l) P_2^1(\cos \theta') \cos \phi' B_{\text{ext}} \\ & - \frac{1}{3}A_{y'z'}^l (\chi_{y'}^l + \chi_{z'}^l) P_2^1(\cos \theta') \sin \phi' B_{\text{ext}} \\ & + \frac{1}{6}(A_{x'x'}^l \chi_{x'}^l - A_{y'y'}^l \chi_{y'}^l) P_2^2(\cos \theta') \cos 2\phi' B_{\text{ext}} \\ & + \frac{1}{6}A_{x'y'}^l (\chi_{x'}^l + \chi_{y'}^l) P_2^2(\cos \theta') \sin 2\phi' B_{\text{ext}}, \end{aligned} \quad (7)$$

where the terms P_l^m represent the usual associated Legendre polynomials [24].

Equation (7) can be simplified by noting that for a cubic system the magnetic susceptibility is isotropic and here defined as χ . We also have $\theta' = 90^\circ$, as of course our field is per definition perpendicular to the rotation axis and ϕ' represents

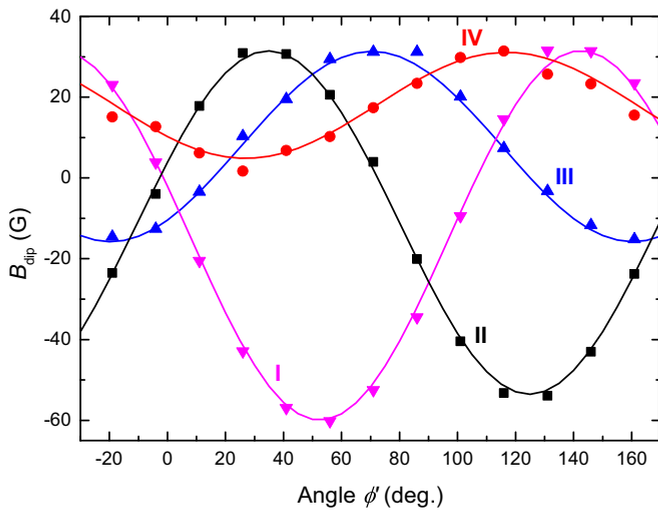


FIG. 5. (Color online) Angular dependence of the dipolar contributions of the four frequencies reported on Fig. 4. The symbols are obtained after the subtraction described in Eq. (3). The lines correspond to the theoretical calculations positioning the muon at the site $(0.532, 0.532, 0.532)$ and taking into account the rotation axis (see text).

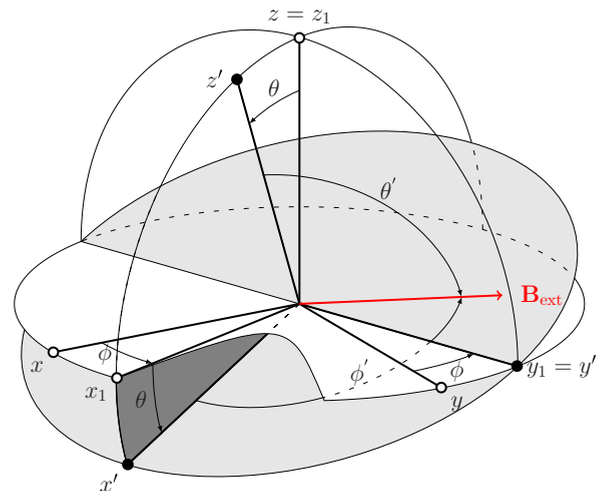


FIG. 6. (Color online) Definition of the Euler angles defining the rotation axis and of the polar and azimuth angles defining the direction of the external field. The reference frame (x, y, z) is the reference frame of the crystal and (x', y', z') is the one defined by the rotating plane and rotation axis.

TABLE II. Fit results obtained adjusting Eq. (8) on the angular dependence of the dipolar contribution for each signal.

	$A_{x'x'}^l$ (mol/emu)	$A_{y'y'}^l$ (mol/emu)	$A_{x'y'}^l$ (mol/emu)
Signal I	-0.0128(38)	-0.1713(11)	-0.2925(31)
Signal II	0.0235(30)	-0.1662(32)	0.2617(26)
Signal III	-0.0640(38)	0.1747(35)	0.0945(27)
Signal IV	0.0538(48)	0.1627(51)	-0.0637(41)

our rotation angle. We therefore have

$$\begin{aligned}
 B_{\text{dip},\parallel} = & \frac{1}{2} \chi (A_{x'x'}^l + A_{y'y'}^l) B_{\text{ext}} \\
 & + \frac{1}{2} \chi (A_{x'x'}^l - A_{y'y'}^l) \cos 2\phi' B_{\text{ext}} \\
 & + \chi A_{x'y'}^l \sin 2\phi' B_{\text{ext}}. \quad (8)
 \end{aligned}$$

Note that the local field at the muon site $\mathbf{B}_{\text{loc,TF}}$ is not exactly parallel to \mathbf{B}_{ext} as in general the dipolar tensor is not diagonal. However, as $|\mathbf{B}_{\text{ext}} + \mathbf{B}_{\text{cont}} + \mathbf{B}_{\text{Lor}} + \mathbf{B}_{\text{dem}}| \gg |\mathbf{B}_{\text{dip}}|$, it is enough to consider solely the component of \mathbf{B}_{dip} along the direction of the external field [the differences are here of the order of 0.1% on the values of B_{dip} obtained by Eq. (3)].

As B_{ext} (5200 G) and χ (0.030 emu/mol [25–27]) are known, the task is now to obtain the elements $A_{x'x'}^l$, $A_{y'y'}^l$, and $A_{x'y'}^l$ by fitting the angular dependence of each muon frequency [28]. The results of the fits obtained by adjusting Eq. (8) on the angular dependence of each signal are shown in Table II.

At this point, these results can be used to solve Eq. (6) for each signal (site) and obtain the Euler angles θ and ϕ of the rotation axis, as well as the parameter a_{dip} characterizing the representation of the dipolar tensors \mathbf{A}_{dip} in the reference frame of the crystal (see Table I). As each signal should provide us with the same parameters, we have a rather precise determination of these parameters, i.e., $a_{\text{dip}} = -0.2044(40)$ mol/emu, $\theta = 83(1)^\circ$, and $\phi = 242(1)^\circ$. We note that the values of θ and ϕ could be confirmed *a posteriori* by performing x-ray Laue measurements of the crystal. But we stress that the values of θ and ϕ extracted by μ SR must be considered as more precise as, for example, they take into account any possible misalignment of the sample mounting compared to the direction of the external magnetic field. The experimentally determined value of a_{dip} can be directly compared with dipolar sum calculations. Figure 7, which exhibits the theoretical value of a_{dip} calculated for different sites $4a$, indicates that this experimental value is compatible with muons sitting either at the site (0.532,0.532,0.532) or (0.721,0.721,0.721). We stress that a small uncertainty on the magnetic susceptibility (and therefore finally on the obtained value of a_{dip}) will only weakly affect the determination of the muon site (for example, an uncertainty as high as $\pm 10\%$ on χ leads to a shift of $\Delta x = \pm 0.005$ on the muon-site coordinates). The obtained sites correspond to almost symmetrical positions along the crystal diagonal on each side of the plane formed by three adjacent Mn ions (see Fig. 8). However, the site (0.721,0.721,0.721) appears not probable as located rather close to the Si ion on the crystal diagonal ($r_{\mu\text{-Si}} \simeq 1.0 \text{ \AA}$). Note that such a short μ -ion distance can sometimes be observed

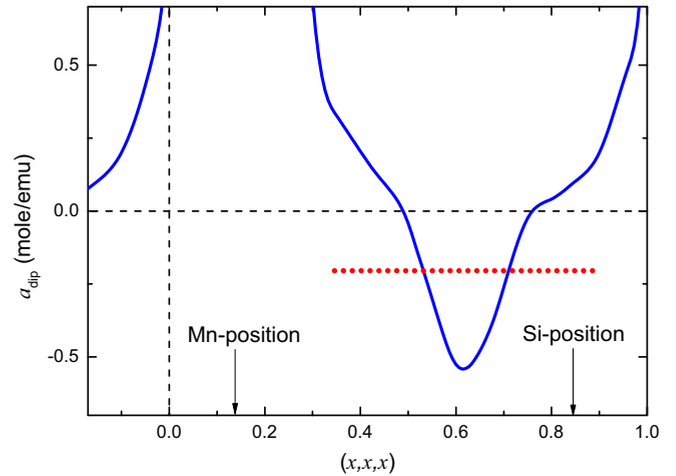


FIG. 7. (Color online) Dipolar sum calculation of the parameter a_{dip} characterizing the representation $[A_{ij}^c]$ of the dipolar tensors in the crystal reference frame for the $4a$ sites. The divergence occurs at the $4a$ position of the Mn ion. The red dotted line corresponds to the results of the fits as explained in the text.

when a bounding between the muon and a negative ion occurs, as in the high- T_c cuprates.

To demonstrate the perfect agreement between theoretical calculations and experiment, the lines in Fig. 5 represent the dipolar calculations for the site (0.532,0.532,0.532) with the Euler angles $\theta = 83^\circ$ and $\phi = 242^\circ$. Note that as the four sites are crystallographically equivalent, one expects similar amplitudes of the four signals, as experimentally observed. We note also that our measurements performed with angles of $\phi' = 86^\circ$ and -4° correspond, respectively, to an external field applied almost into a principal plane [i.e., (110)] and almost along the principal axis $[00\bar{1}]$ (as the Euler angle for

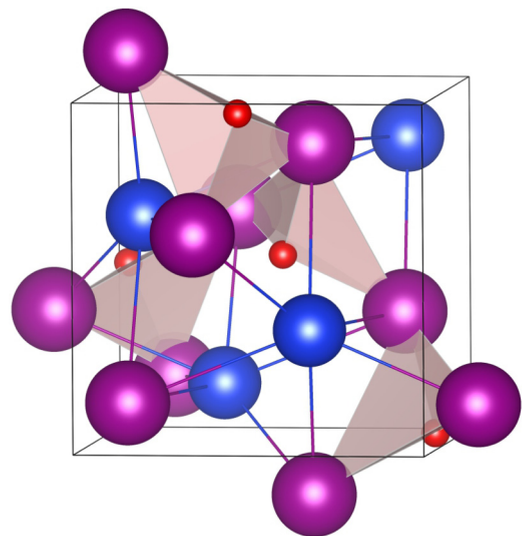


FIG. 8. (Color online) Sketch of the crystallographic structure of MnSi (Mn ions are drawn in purple, Si ions in blue). The muon position (0.532,0.532,0.532) is also indicated (red) as well as the other three equivalent sites. Note that six Mn ions, which do not belong to the primary unit cell, are also displayed.

the rotation axis is $\theta = 83^\circ$). Figure 3 shows that the signals tend to collapse into two pairs in the first case and all together (and with a dipolar contribution tending to zero) in the second case, as expected. For completeness, we would also like to mention the possibility of a small lattice dilation around the muon. We notice first that such effects are usually extremely small in metals (of the order of the percent, if detectable at all; see, for example, Ref. [29]). A hypothetical weak dilation of the lattice would decrease the calculated absolute values of the a_{dip} parameter, which would push slightly the muon position in the direction of the plane formed by the three adjacent Mn ions (see Fig. 8). However, the main conclusion that the muon is stopped at a $4a$ position would not be affected.

Finally, we note that a byproduct of our analysis is the determination of the contact field \mathbf{B}_{cont} . Basically two interactions contribute to the contact field. The first one is a result of the Pauli paramagnetism of the conduction electrons and their Fermi contact interaction with the muon [21]. In systems with localized moments, as MnSi, an additional contribution arises as the spin polarization of the conduction electrons at the muon site will be increased through the Ruderman-Kittel-Kasuya-Yosida interaction due to the local magnetic moments. As the Pauli susceptibility of the conduction electrons is much smaller than the magnetic susceptibility from the local $3d$ moments (i.e., $\chi \simeq \chi_{3d} \gg \chi_{\text{Pauli}}$), the Fermi interaction can be safely neglected in the contact field, which therefore can be written as

$$\mathbf{B}_{\text{cont}} = \mathbf{A}_{\text{cont,TF}} \chi \mathbf{B}_{\text{ext}}, \quad (9)$$

where \mathbf{A}_{cont} is the hyperfine contact coupling tensor. By writing Eq. (2), we note that we implicitly assumed that the contact field is independent of the direction of \mathbf{B}_{ext} (i.e., $\mathbf{A}_{\text{cont,TF}} = A_{\text{cont,TF}} \mathbf{E}$) as observed in a number of compounds with local moments (see, for example, Ref. [31] and references therein). In any case, even if one assumes that the weak angular dependence of $\bar{\nu}$ is solely arising from a hypothetical angular dependence of \mathbf{A}_{cont} , this would correspond to a variation of less than 4% of the hyperfine contact coupling tensor during the rotation. As we *know* that some angular dependence has to arise from the demagnetization factor (see Fig. 1), this number has to be considered as an upper limit. By assuming that the hyperfine contact coupling tensor is isotropic, and taking into account that for our sample geometry the average value of the demagnetization factor is $\bar{N} = 0.47 \times (4\pi)$ one obtains a value of $A_{\text{cont,TF}} = -0.9276(20)$ mol/emu, which can now be included in the computation of the expected values of the spontaneous fields occurring at the muon sites below T_C as described in the next section.

B. Zero-field data discussion

At this point, having a solid knowledge of the muon stopping site, it appears legitimate to thoroughly discuss the number and values of the spontaneous μ frequencies observed below T_C [14,16,17] in zero-applied field (ZF) μ SR experiments. All the μ SR studies reported to date reveal the occurrence of two spontaneous μ frequencies with values $\nu_{\text{ZF},1} \simeq 12.3$ MHz and $\nu_{\text{ZF},2} \simeq 28.0$ MHz for $T \rightarrow 0$ K. Our first task here will be to discuss why solely two frequencies are observed.

The magnetic structure of MnSi is characterized by spins forming a left-handed incommensurate helix with a propagation vector $k \simeq 0.036 \text{ \AA}^{-1}$ in the [111] direction [5–7]. The static Mn moments ($\sim 0.4\mu_B$ for $T \rightarrow 0$ K) point in a plane perpendicular to the propagation vector. The period, which is incommensurate to the lattice constant, is about 18 nm. Due to the incommensurability of the magnetic structure, one expects a continuous set of local fields at our $4a$ sites, and therefore the field distribution $D_i(B_{\text{loc,ZF}})$ at each site must be considered (here the index i distinguishes the four subsites $4a$). It was shown that such a magnetic structure leads to a field distribution given by [21,32,33]

$$D_i(B_{\text{loc,ZF}}) = \frac{2}{\pi} \frac{B_{\text{loc,ZF}}}{\sqrt{B_{\text{loc,ZF}}^2 - B_{\text{min},i}^2} \sqrt{B_{\text{max},i}^2 - B_{\text{loc,ZF}}^2}}, \quad (10)$$

and is characterized by two peaks due to the minimum and maximum cutoff field values. Hence, at first glance, one would expect to observe up to eight peaks in the Fourier spectra of the ZF- μ SR data. However, as the sites *II*, *III*, and *IV* are located symmetrically around the direction [111] of the propagation vector (which represents for them a threefold symmetry axis), it can be shown that muons stopping at these sites will sense identical field distributions, given by Eq. (10), i.e., $D_{II}(B_{\text{loc,ZF}}) = D_{III}(B_{\text{loc,ZF}}) = D_{IV}(B_{\text{loc,ZF}})$. On the other hand, muons stopping at the site *I* will all sense a unique field value irrespective of the phase of the helix at the muon stopping site. Hence, purely geometrical considerations already reduce the maximum number of peaks in the Fourier spectra of the ZF- μ SR data down to three. Note that this conclusion is not affected if the sample is not monodomain. Hence for the other possible magnetic domains (which, by taking into account the helicity, are characterized by propagation vectors along $[\bar{1}\bar{1}1]$, $[\bar{1}1\bar{1}]$, and $[1\bar{1}\bar{1}]$) one obtains the same field distributions as for the original domain; albeit the symmetry of the sites with respect to the magnetic structure will be accordingly permuted.

To gain more insight, we have computed the expected theoretical cutoff field values of the field distributions. For a given type of muon site, the spontaneous local field in the incommensurate magnetic phase is given by [30]

$$\mathbf{B}_{\text{loc,ZF}}(\mathbf{r}'_i) = \mathbf{B}_{\text{dip,ZF}}(\mathbf{r}'_i) + \mathbf{B}_{\text{cont,ZF}}(\mathbf{r}'_i). \quad (11)$$

As before, the index i distinguishes the four subsites $4a$ and runs from *I* to *IV*. Since the magnetic structure of MnSi is incommensurate, the spontaneous field differs from one particular site of type i to another site of the same type. In Eq. (11) \mathbf{r}'_i represents the set of vectors defining the positions of the sites of type i with $\mathbf{r}'_i = \mathbf{r}_i + \mathbf{R}$, where \mathbf{r}_i defines the position of the site of type i in the primitive cell and \mathbf{R} is a vector belonging to the direct crystal lattice. The calculation of the dipolar contribution on the right-hand side of Eq. (11) is straightforward and will fully be determined by the knowledge of the muon stopping site and the details of the magnetic structure, such as propagation vector, value, and direction of the magnetic moments, as well as helicity of the helix. On the other hand, the contact contribution can be written as

$$\mathbf{B}_{\text{cont,ZF}}(\mathbf{r}'_i) = V_{\text{mol}} A_{\text{cont,ZF}} \mathbf{M}(\mathbf{r}'_i), \quad (12)$$

where V_{mol} is the volume of one mol of Mn-ions. In writing Eq. (12), we have assumed that the contact coupling is isotropic

as suggested from our TF data. $\mathbf{M}(\mathbf{r}'_i)$ is the *local* magnetization at the muon site \mathbf{r}'_i . To compute $\mathbf{B}_{\text{cont,ZF}}$, the first natural choice is to set $A_{\text{cont,ZF}}$ equal to $A_{\text{cont,TF}}$ determined above. This is reasonable assuming that no massive changes occur on the Fermi surface when crossing T_C and considering that due to the long wavelength, the close Mn neighbors around the muon, which are determining the strength of the RKKY interaction, are essentially ferromagnetically aligned in the magnetic phase. This is the only choice made for the calculation, as the *local* magnetization $\mathbf{M}(\mathbf{r}'_i)$ is also fully determined by the details of the magnetic structure and is given by

$$\mathbf{M}(\mathbf{r}'_i) = \frac{1}{V_{\text{nn}}} \sum_{j=1}^N \mathbf{m}_j = \frac{4}{NV_{\text{cell}}} \sum_{j=1}^N \mathbf{m}_j, \quad (13)$$

where the volume $V_{\text{nn}} = (NV_{\text{cell}})/4$ is the volume occupied by the N Mn ions involved in the sum. The direction of the Mn moments \mathbf{m}_j is determined by the details of the magnetic structure. For our calculations, as the RKKY interaction is local, we have restricted the sum to a sphere having a radius of one lattice constant around the muon site.

The values of $\mathbf{B}_{\text{loc,ZF}}(\mathbf{r}'_i)$ are located on the path of an ellipse for which the semimajor and semiminor axis values are B_{max} and B_{min} . Starting from a known magnetic structure, the details for a precise computation of these values is thoroughly explained in Ref. [21]. The present calculations were performed on a sphere containing more than 5×10^5 unit cells. By setting $A_{\text{cont,ZF}} = A_{\text{cont,TF}}$, we stress again that our calculation does not contain any free parameters as the magnetic structure is taken from the literature, the muon site is determined by our TF data and the contact coupling constant is taken *as is* from the TF data. From this calculation, one obtains $B_{\text{max},II,III,IV} = 2065$ G (corresponding to a muon frequency of 28.0 MHz), $B_{\text{min},II,III,IV} = 900$ G (12.2 MHz), and $B_I = 880$ G (11.9 MHz). The lower panel of Fig. 11 shows the obtained field distributions which have been convoluted for better visibility (see caption). The red contribution represents the sum of the field distributions for the *II*, *III*, and *IV* sites whereas the blue contribution shows the field distribution expected at the *I* site.

This calculation can now be directly compared to the analysis of a high-statistics ZF- μ SR measurement performed at 5 K on a single crystal obtained according to the same procedure as the one used for the TF- μ SR measurements. We note first that the oscillatory part of the muon polarization function associated with the field distribution given by Eq. (10), and given by

$$P_{\text{osc},i}(t) = \int_{B_{\text{min},i}}^{B_{\text{max},i}} D_i(B_{\text{loc,ZF}}) \cos(\gamma_\mu B_{\text{loc,ZF}} t), \quad (14)$$

cannot be obtained analytically. We therefore approximate $D_i(B_{\text{loc,ZF}})$ with a shifted Overhauser distribution,

$$D_i(B_{\text{loc,ZF}}) \simeq \frac{1}{\pi} \frac{1}{\sqrt{\Delta B_i^2 - (B_{\text{loc,ZF}} - B_{\text{av},i})^2}}, \quad (15)$$

where $\Delta B_i = (B_{\text{max},i} - B_{\text{min},i})/2$ and $B_{\text{av},i} = (B_{\text{max},i} + B_{\text{min},i})/2$. The upper panel of Fig. 9 shows the difference between the exact and approximated field distribution, which is symmetrical with respect to the singularities, i.e., some weight is transferred from the upper cutoff to the lower cutoff

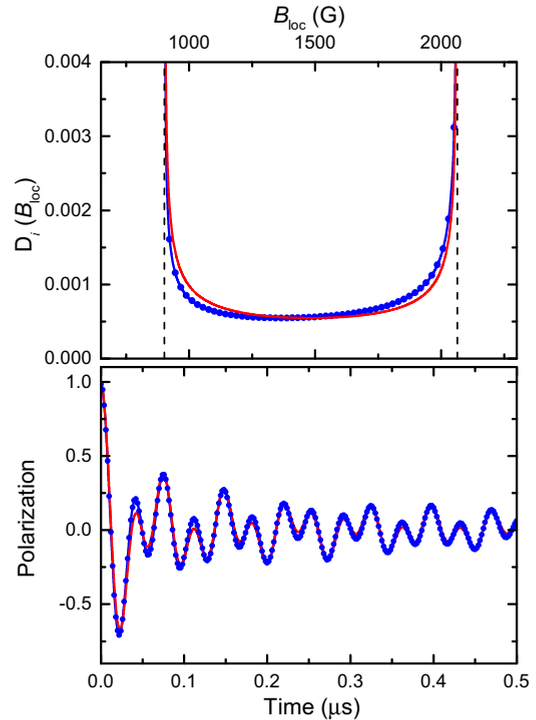


FIG. 9. (Color online) (Upper panel) Exact [blue line and symbols; see Eq. (10)] and approximated [red line; see Eq. (15)] field distributions expected at sites *II*, *III*, and *IV*. (Lower panel) Muon polarization function deduced from the field distributions shown on the upper panel (see also text).

field. The lower panel of Fig. 9 shows the oscillatory part of the muon polarization function obtained for both field distributions. For the exact field distribution a numerical calculation was performed using Eq. (14), whereas for the approximated field distribution the muon polarization function can be obtained analytically and is given by

$$P_{\text{osc Overh},i}(t) = J_0(\gamma_\mu \Delta B_i t) \cos(\gamma_\mu B_{\text{av},i} t), \quad (16)$$

where J_0 is a Bessel function of the first kind. We see that the function $P_{\text{osc Overh},i}(t)$ catches the essential features of $P_{\text{osc},i}(t)$ and constitutes a good approximation. Having this in mind, we can now write a function which will be fitted to the ZF experimental data, i.e.,

$$\begin{aligned} A P_{\text{osc ZF}}(t) = & \mathcal{A}_1 \cos(\gamma_\mu B_1 t + \psi_1) \exp(-\lambda_1 t) \\ & + \mathcal{A}_2 J_0(\gamma_\mu \Delta B_2 t) \cos(\gamma_\mu B_{\text{av},2} t + \psi_2) \\ & \times \exp(-\lambda_2 t), \end{aligned} \quad (17)$$

where the first component is associated with site *I* and the second with sites *II*, *III*, and *IV*. The depolarization rates reflect as usual any static or dynamical effects, the discussion of which goes beyond the scope of this study. Figure 10 shows the ZF- μ SR spectra with the fitted polarization function. From the fitted parameters one obtains $B_I^{\text{exp}} = 911(2)$ G (12.34 MHz), $B_{\text{min},II,III,IV}^{\text{exp}} = 959(3)$ G (12.99 MHz), and $B_{\text{max},II,III,IV}^{\text{exp}} = 2071(3)$ G (28.06 MHz). These values agree very well with our calculations performed without free parameters. In addition, both phase parameters ψ_1 and ψ_2 were found to be compatible with zero, as expected. The depolarization rates are found to

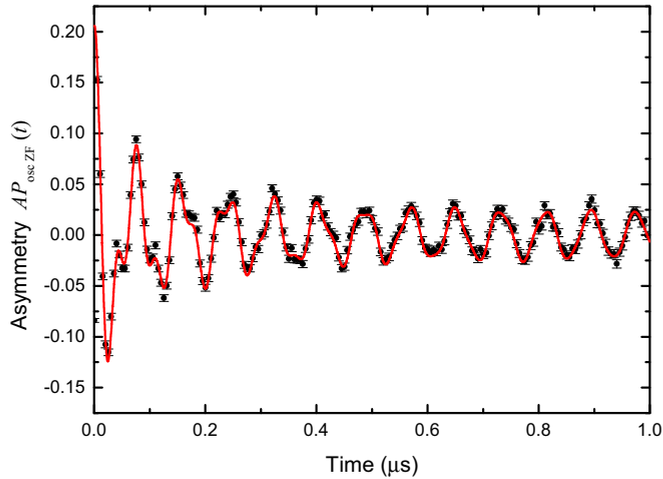


FIG. 10. (Color online) ZF- μ SR data taken at 5 K (initial muon polarization parallel to the [111] direction). The line represents a fit of Eq. (17) to the data using the parameters given in the text (for better visibility a time binning corresponding to 5 ns was chosen, whereas the fits were performed with a time binning of 1.25 ns).

have the values $\lambda_1 = 0.70(03) \mu\text{s}^{-1}$ and $\lambda_2 = 1.78(16) \mu\text{s}^{-1}$. We note that the small difference between the values of B_I^{exp} and $B_{\text{min},II,III,IV}^{\text{exp}}$ (which is also reflected in our calculations) perfectly explains that solely two peaks were invoked in former μ SR data realized with much less statistics. Finally an interesting point is the fact that $B_I < B_{\text{min},II,III,IV}$ in both the calculations and the experimental data. This is remarkable as the position of $B_{\text{min},II,III,IV}$, with respect to B_I , depends on the helicity of the incommensurate helix and is located above B_I only for left-handed helicity. It is therefore tempting to take the ZF- μ SR data as a late confirmation of the helix helicity. We note also that a close look at the fast Fourier transform reveals as expected a very slight divergence near the cutoff fields between the data and the shifted Overhauser distribution (see the upper panel of Fig. 11 and compare the data points to the red distribution). We therefore conclude here that from the knowledge of the muon site determined by our TF data, one can utterly explain the characteristics of the ZF- μ SR data.

C. Muon site: comparing experimental determination and *ab initio* calculations

For completeness, we compare now our precise experimental determination of the muon site with *ab initio* calculations based on the density-functional theory (DFT). Such calculations have recently been shown to accurately reproduce the muon sites observed in different materials as wide-gap semiconductors, insulating systems or cuprates and iron-based high- T_c superconductors [34–37]. The calculations were performed at the University of Parma and we like to stress that they have been performed in a *blind modus*, i.e., without any prior knowledge of the muon site deduced from the TF- μ SR data.

To analyze the electronic structure of MnSi we use the FP-LAPW approach as implemented in the ELK code [38]. The generalized gradient approximation (GGA) as formulated by Perdew, Burke, and Ernzerhof is used to approximate the exchange and correlation potential [39]. The cutoff for

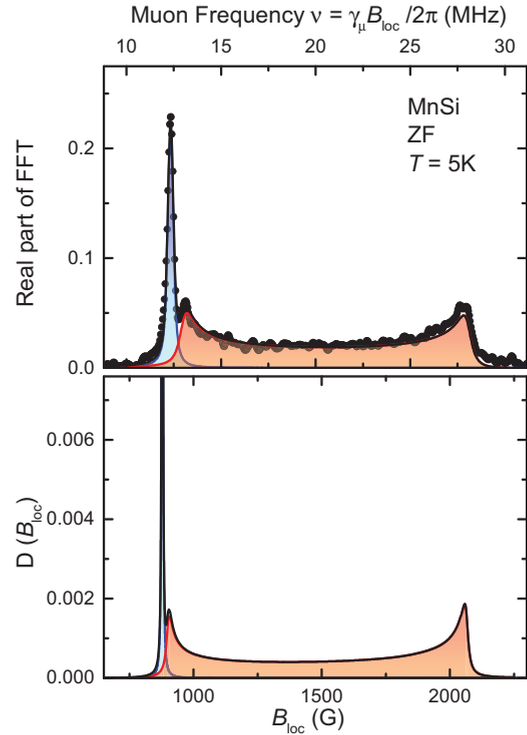


FIG. 11. (Color online) (Upper panel) Fast Fourier transform of the ZF- μ SR data obtained at 5 K and shown on Fig. 10. The blue and red components represent, respectively, the first and second component of the right-hand side of Eq. (17). (Lower panel) Computed field distributions for the I site (blue) and for the II , III and IV sites (red). For better visibility, these field distributions have been slightly folded by Lorentzian distributions with widths (full width at half maximum) of 5 and 20 G, respectively.

the plane wave expansion in the interstitial region is set to $|\mathbf{G} + \mathbf{k}|_{\text{max}} \cdot R_{\text{mt}} = 9$ (R_{mt} being the smallest muffin-tin radius). The reciprocal space is sampled on a $16 \times 16 \times 16$ Monkhorst-Pack [40] grid. The experimental atomic positions and lattice constant are used. A ferromagnetic ground state is considered and the resulting magnetic moment on the Mn atoms is approximately $1 \mu_B$. Indeed MnSi in a well-known case in which the mean-field approximation leads to an overestimation of the magnetic moments [41,42].

The muon site is identified from the minima of the electrostatic potential obtained from the ground-state electronic density of the unperturbed material. There are four equivalent minima in the unit cell, as shown in Fig. 12, which correspond to the $4a$ Wyckoff position with fractional coordinates (0.523, 0.523, 0.523). This coarse estimation nicely agrees with the experimentally evaluated site.

The standard procedure [34,36] for the DFT site assignment further requires the analysis of the perturbation introduced by the muon in the vicinity of its embedding site. This is done by analyzing the electronic and crystallographic modifications introduced by the charged impurity. The muon site is eventually validated by considering the spread of its wave function. These steps are computationally demanding and outside the scope of the present work. However preliminary indications suggest that the inclusion of the muon in a supercell produces an even

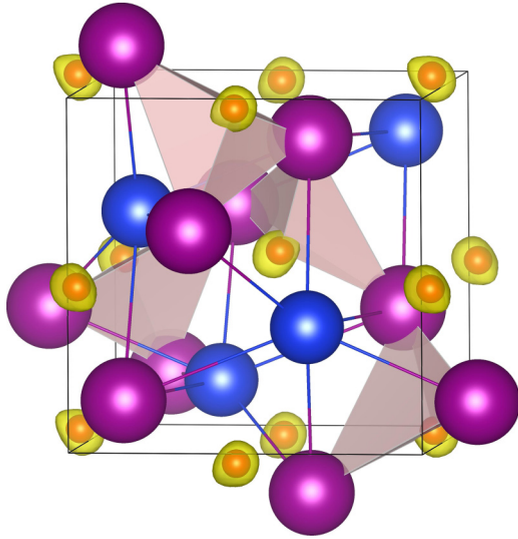


FIG. 12. (Color online) Sketch of the crystallographic structure of MnSi with minima regions of the electrostatic potential (yellow). The rest of the color scheme is the same as in Fig. 8. The yellow regions define isosurfaces with an energy set to $V = E_0/2$ with E_0 being the ground-state energy obtained from the solution of the Schrödinger equation for the muon in the electrostatic potential. The equivalent muon stopping sites determined by TF- μ SR are also indicated and are enclosed in the minima regions. Note that compared to Fig. 8, we also report the muon sites located just outside of the primary unit cell, which are enclosed in minima regions extending into the primary unit cell.

closer agreement with the experimental value, providing a $4a$ muon stopping site at the coordinates (0.538,0.538,0.538).

IV. CONCLUSIONS

We conclude that the observed angular dependence of the TF- μ SR signals (except for a very slight dependence due to the demagnetization factor) can be beyond any doubt ascribed by a muon sitting at a $4a$ Wyckoff position and detecting the dipolar field produced by the Mn moments. Consequently the μ SR response of paramagnetic MnSi under the application of a magnetic field can be fully understood without invoking a hypothetical magnetic polaron state. In addition we have shown that the knowledge of the muon stopping site provides a clear understanding of the ZF- μ SR data and that the computed field distribution perfectly agrees with the measured one. Moreover, *blind modus ab initio* DFT calculations provide a $4a$ muon stopping which is in very close agreement with the site determined by the experimental μ SR data.

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