Intrinsic paramagnetism and aggregation of manganese dopants in SrTiO$_3$

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Using local-probe magnetic-characterization techniques of muon spin relaxation and electron spin resonance we have investigated the Mn-induced magnetism of the wide-band-gap perovskite SrTiO$_3$. Our results clearly demonstrate that this diluted magnetic oxide remains paramagnetic down to low temperatures for both doping cases, i.e., when Mn substitutes for Sr or Ti. In addition, both experimental techniques have revealed that the distribution of individual Mn$^{2+}$ and Mn$^{4+}$ ions is nonrandom, as these ions partially aggregate into nanosized clusters.

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I. INTRODUCTION

Oxide semiconductors with a fraction of the host cations substituted by magnetic ions, i.e., diluted magnetic oxides (DMOs), have been presently intensively investigated for potential spintronic applications [1–8]. The quest for new multiferroic/magnetoelectric DMO materials that would be of great technological importance is, however, hampered by the general concern whether their magnetism, coupled or uncoupled to a polar state of the material, is undoubtedly intrinsic [9–12]. Lately, there has been huge interest in the electronically and chemically doped wide-band-gap perovskite SrTiO$_3$. In this incipient ferroelectric that is highly polarizable, yet it misses the electric freezing even at $T \to 0$ due to quantum fluctuations [13], a small concentration of dopants can drastically change its electric and magnetic properties. To name only the most prominent examples, superconductivity [14,15], a magnetoelectric multiglass state [16,17], a two-dimensional electron gas [18,19], and ferromagnetism [20–22] have been reported in the bulk or at the surface of SrTiO$_3$ and at interfaces with other insulating oxides.

Manganese impurities in SrTiO$_3$, in particular, have been thoroughly investigated because they can be multivalent and because of their polar and magnetic moments. They are held responsible for a relaxor-type polar state [23] when Mn$^{2+}$ substitutes for Sr$^{2+}$ (at the dodecahedral A site), while on the other hand, the material remains quantum paraelectric when Mn$^{4+}$ substitutes for Ti$^{4+}$ (at the octahedral B site) [24,25]. The polar instability observed in the case of the A-site doping is due to off-center location of the much smaller Mn$^{2+}$ ion with respect to the Sr$^{2+}$ ion and was confirmed by experiments [26–28] and theory [29]. The polar freezing has been further argued to induce a spin-glass state for certain compositions, where a magnetoelectric coupling should lead to a novel magnetoelectric multiglass [16,17]. The existence of such a state has been recently hotly debated [30–32]. As an alternative, the frozen magnetism was suggested to be due to segregation of magnetic species into Mn$_3$O$_7$-like clusters and thus to be extrinsic [25,30]. Moreover, such magnetism was shown to critically depend on synthesis conditions [30]. Thus, the intrinsic magnetic ground state of the Mn-doped SrTiO$_3$ remains elusive and calls for an in-depth magnetic study. Furthermore, the role and size of superexchange interactions between dopants [33] in SrTiO$_3$ is unclear. These magnetic interactions could further be influenced by a seemingly inevitable presence of oxygen vacancies (V$_O$) in these materials [34], which might lead to bound magnetic polarons [35]. This mechanism could then explain the ferromagnetic ordering of some Mn-doped SrTiO$_3$ materials [34], but needs to be critically assessed.

In order to address all the above-mentioned pending issues about the Mn-doped SrTiO$_3$, we have performed an in-depth local-probe investigation. Here, we combine results of muon spin relaxation ($\mu$SR) and electron spin resonance (ESR) investigations. A further insight into the dopant-induced electronic states in SrTiO$_3$ has been provided by diffuse reflectance spectroscopy (DRS) measurement. We show that all the investigated samples, including the A-site-doped ones, remain paramagnetic down to low temperatures and that the Mn ions tend to form dopant-reach islands. Similar nonrandom distributions of dopants on a nanoscale, spinodal decomposition into regions with high/low concentration of dopants and the same crystal structure as the semiconducting host, were observed before in different diluted magnetic semiconductors [5,36–39]. However, in doped SrTiO$_3$, such aggregation has not been observed yet and is fundamentally different from the recently reported [40] tendency of the Mn$^{2+}$ ions to segregate in grain boundaries.

II. EXPERIMENTAL DETAILS

Powder samples were synthesized according to the synthesis protocols thoroughly explained in Ref. [30]. Our study benefits from a systematic synthesis approach employing high-energy homogenization of starting reagents and various different sintering conditions (oxygen partial pressures were varied by applying either O$_2$ or N$_2$ atmosphere in post treatment), thus allowing a production of high-quality powder samples of the cubic perovskite SrTiO$_3$, nominally doped on
both A and B sites in the doping range from 1% to 3%. For each doping concentration and a chosen doping site, the O2 and N2 post-treated samples were obtained from the same parent compound. X-ray diffraction patterns showed no indication of secondary phases in any of the investigated samples. The 3%-doped samples used in our study were the same as in Ref. [30]. In the following, our samples are labeled by XcY, where X = A,B denotes the two sites, c the doping concentration in percent, and Y = O2,N2 the post-treatment atmosphere; e.g., A3O2 stands for the 3% nominally A-site Mn-doped sample that was post-treated in the O2 atmosphere.

Magnetism on a local scale was inspected with μSR and ESR. The μSR data were collected on the General Purpose Surface-muon (GPS) instrument at the Swiss Muon Source (SμS), Paul Scherrer Institute (PSI), Switzerland. The measurements were performed in the weak transverse field (wTF) of 3 mT, in zero field (ZF), and in various longitudinal fields (LF) applied along the initial muon polarization in the temperature range 1.6–120 K. The ESR measurements were performed between 5 and 300 K in the X band (at 9.5 GHz) on a home-built spectrometer. The DRS was performed at room temperature on the Lambda 650 UV-vis spectrophotometer (Perkin-Elmer) equipped with a diffusion sphere.

III. RESULTS

A. Muon spin relaxation

To assess the reportedly intrinsic presence of the frozen spin-glass state in the A-site-doped SrTiO3 samples [16,17] on a local scale, we first resort to μSR, a technique well known for its extreme sensitivity to local magnetic fields [41]. The muons, that are initially almost 100% spin polarized, stop in the sample and experience a local magnetic field Bμ, which leads to a time-dependent muon polarization P(t). Its form reflects a magnitude, distribution, and fluctuations of Bμ and depends of the magnetic coupling of the muon with its surroundings. If the field is rapidly fluctuating v ≥ γμBμ, where v denotes the field fluctuation frequency and γμ = 2π × 135.5 MHz/T is the muon gyromagnetic ratio, P(t) will decay monotonically with time in ZF. A quasistatic local field, on the other hand, causes a coherent oscillation of the ZF muon polarization that is damped by slow muon dynamics (v ≪ γμBμ) and by a field distribution. The resulting depolarization remains nonmonotonic even for randomly oriented frozen dense (e.g., pertinent to nuclear magnetism [42]) or diluted magnetic moments (e.g., arising from a dilute-alloy spin glass [43]).

In Fig. 1, the μSR asymmetry G(t) (spatial asymmetry of muon decay detected by emitted positrons) that is proportional to P(t) is plotted for the A3O2 sample. The wTF measurements [Fig. 1(a)] are generally accepted as an effective tool to trace the volume fraction of the frozen spins, as the internal fields are typically well above the applied field and thus cause a loss of the precession amplitude. In diluted spin glasses though, the average internal field may be of the order of only 0.1–1 mT, i.e., much smaller than the applied wTF. However, the spin-glass transition should still manifest in a sharp increase of the muon depolarization rate below the transition temperature because the onset of random internal fields broadens the distribution of local fields [44]. Experimentally, we observe a smooth temperature dependence of the transverse relaxation rate λt in the wTF experiment (vide infra), with no notable irregularities (Fig. 2) at the reported spin-glass transition of around 40 K [16,17,31]. The observed damping of the wTF signal can in principle be either due to dephasing in an inhomogeneous static field or due to spin dynamics. These two effects can usually be separated in the ZF experiment. In ZF, the dynamical fields in a diluted magnetic system lead to the monotonic “root-exponential” relaxation [43]

\[ G_{ZF}(t,v) = G_0 e^{-\sqrt{4\pi v/\nu}} = G_0 e^{-2\pi \nu}, \quad (1) \]

where G0 denotes the initial asymmetry and a/γμ the field-distribution width. Static internal fields, on the other hand, in spin glasses lead to a nonmonotonic single-dip relaxation function

\[ G_{LKT}^{LKT}(t) = G_0 \left[ \frac{1}{2} + \frac{a}{2} (1 - at)e^{-at} \right], \quad (2) \]

known as the Lorentzian Kubo-Toyabe (LKT) function [43]. In the A3O2 sample, both models give a similar quality of fit at 1.6 K [see ZF data in Fig. 1(c)], with either the dynamical longitudinal relaxation rate λl = 0.30(1) s⁻¹ [model (1)] or the static field-distribution width a/γμ = 0.32(2) mT [model (2)]. We note that in the latter case the dip in G_{ZF}^{LKT}(t) falls outside

FIG. 1. (Color online) The temperature-dependent μSR asymmetry of the A3O2 sample in (a) wTF and (b) ZF. (c) Field decoupling of the longitudinal μSR relaxation in various LFs at 1.6 K for the same sample. The solid lines in panels (a), (b), and (c) correspond to fits to the models (4), (1), and (3), respectively. The dashed line in (b) shows the expected position of the “1/3 tail,” and in (c) the prediction of the static Lorentzian Kubo-Toyabe model.
the experimental time window, therefore, its existence and the leveling of the data at \( G_0 \) at longer times that is characteristic for powder samples [41] cannot be verified.

In order to distinguish between the two scenarios, we turn to the LF experiment. In the static case, already the applied field of the order of a few times \( a/\gamma_\mu \) noticeably narrows the corresponding local-field distribution, which results in a significantly decreased depolarization. For demonstration, we show in Fig. 1(c) the prediction of the static LKT model \([a/\gamma_\mu = 0.32(2) \text{ mT}]\) in the applied field [43] of 10 mT, which should almost completely remove the relaxation. This is evidently inconsistent with the experiment and thus unambiguously proves that local magnetic fields at the muon sites are dynamical. Since magnetic fields from nuclei are generally regarded static on the muon time scale, this finding demonstrates that the ground magnetic state of the A3O2 sample is dynamical.

In the case of dynamical internal fields with a single fluctuation rate \( \nu \), the \( \mu \)SR asymmetry is given in the applied external field \( B_{\text{LF}} \) by [43]

\[
G_{\text{LF}}(t, \nu) = \int_0^\infty G_d(t, \Delta, \nu) \rho(\Delta, a) d\Delta.
\]  

(3)

Here, \( G_d(t, \Delta, \nu) = e^{-\frac{2\Delta_0^2}{\nu^2}} \) is the dynamical relaxation function for a field distribution with the width \( \Delta/\gamma_\mu \) at a given muon site. In a diluted spin system \( \Delta \) is further distributed, which is typically modeled with the Gaussian function \( \rho(\Delta, a) = \sqrt{\frac{2}{\pi a^2}} e^{-a^2/\Delta^2} \) [43]. A simultaneous fit of various LF data sets between 0 and 320 mT [Fig. 1(c)] to the model (3) yields the field-distribution width \( a/\gamma_\mu = 0.75(10) \text{ mT} \) and the fluctuation frequency \( \nu = 9.1(5) \text{ MHz} \) in the A3O2 system at 1.6 K. We note though that the modeling of the decoupling experiment does not perfectly match the experimental data at intermediate fields [see the 40-mT curve in Fig. 1(c)], which suggests that there are regions in the sample with different \( \nu \). For instance, a region with larger \( \nu \) would exhibit faster relaxation than modeled for a single fluctuation rate.

A similar, dynamical behavior was found also in the other two investigated samples: A3N2 and B3O2. The corresponding field-distribution widths and fluctuation frequencies at 1.6 K obtained from \( \mu \)SR relaxation in various LFs are summarized in Table I. Both parameters are found to increase with the increasing occupancy of the B site, despite the fact that the average magnetic-moment size decreases: \( \text{Mn}^{2+} \quad (S = \frac{5}{2}) \) on B sites versus \( \text{Mn}^{2+} \quad (S = \frac{7}{2}) \) on A sites.

The temperature dependence of the relaxation rates is shown in Fig. 2 for all three samples. The longitudinal relaxation rates \( \lambda_l \) were obtained by fitting the ZF data to the “root-exponential” decay [Eq. (1)], while the transverse relaxation rates \( \lambda_t \) correspond to fits of the wTF data to the model

\[
G_{\text{wTF}}(t) = G_0 \cos(\gamma_\mu B_{\text{wTF}} t + \varphi) e^{-\sqrt{2} \varphi t},
\]  

(4)

where \( \varphi \) denotes a small tilt of the initial muon polarization from the beam direction. First, we stress that \( \lambda_l \) and \( \lambda_t \) are nearly equal at each temperature, revealing that the contribution of inhomogeneous static fields to \( \lambda_l \) is marginal. This again confirms the dynamical magnetic state in the A3O2 sample. Second, a minimum of both relaxation rates is observed around 75 K. In the A3N2 sample, the temperature dependence of \( \lambda_l \) is very similar, while in the B3O2 sample this minimum is shallower. The increase of the relaxation rates with decreasing temperature below 75 K is typical of slowing down of spin fluctuations at low temperatures, as \( \lambda = \frac{2\Sigma}{\hbar} \) [43]. Above 75 K, on the other hand, the increase of the relaxation rates is rather unusual.

### Table I. XANES-derived occupancy of the Mn\(^{2+}\) (Mn\(^{4+}\)) ions on the A (B) sites in SrTiO\(_3\) [30], the width of the field distribution at the muon site \( a/\gamma_\mu \) and the field fluctuation frequency \( \nu \) for the three investigated samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>A site (Mn(^{2+})) (%)</th>
<th>B site (Mn(^{4+})) (%)</th>
<th>( a/\gamma_\mu ) (mT)</th>
<th>( \nu ) (MHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A3N(_2)</td>
<td>82</td>
<td>18</td>
<td>0.67(5)</td>
<td>4.4(2)</td>
</tr>
<tr>
<td>A3O(_2)</td>
<td>36</td>
<td>64</td>
<td>0.75(10)</td>
<td>9.1(5)</td>
</tr>
<tr>
<td>B3N(_2)</td>
<td>13</td>
<td>87</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B3O(_2)</td>
<td>9</td>
<td>91</td>
<td>0.85(10)</td>
<td>18.3(4)</td>
</tr>
</tbody>
</table>

### B. Electron spin resonance

In order to obtain further insight into the dynamical magnetic state of the Mn-doped SrTiO\(_3\), we next turn to the ESR measurements. ESR has proven highly informative in determining the Mn-ion valence and its local surroundings in SrTiO\(_3\). The substitution of Mn\(^{4+}\) for Ti\(^{4+}\) on the B site was observed by this technique long ago [45], while the successful Mn\(^{2+}\) for Sr\(^{2+}\) substitution on the A site has been claimed only recently [27]. At high temperatures, both cases are characterized by distinctive hyperfine-split ESR sextets, with the \( g \) factor \( g = 1.994 (2.004) \) and the hyperfine coupling \( a = 7.5 \text{ mT} \) (8.8 mT) for Mn\(^{4+}\) (Mn\(^{2+}\)) [45,46]. Such sextets are consistent only with a cubic local crystal-field symmetry and negligible exchange interactions. For the off-center position of the Mn\(^{2+}\) ions at the A sites, the absence of...
we observe any Mn$^{3+}$ (see Table II). We stress that in neither of our samples did $\chi_b$ bulk susceptibilities significantly reduce a of the absorption line or from the Lorentzian distribution of 3%-doped samples are compared in Fig. 3. They can be nicely the excess charge of vacancies. The room-temperature $X$-band ESR spectra of all four 3%-doped samples are compared in Fig. 3. They can be nicely fit to a sum of two Lorentzian-broadened sextets (components C1 and C2) and a much broader C3 component that also exhibits the Lorentzian shape. We note that the Lorentzian shape can either result from motional/exchange narrowing [48] of the absorption line or from the Lorentzian distribution of the local fields in a random dilute system due to $1/r^6$-type interactions [49]. The position and the splitting of the observed sextets are characteristic of the Mn$^{2+}$ ions at the A site (C1 sextet) and the Mn$^{4+}$ ions at the B site (C2 sextet) [45,46] (see Table II). We stress that in neither of our samples did we observe any Mn$^{3+}$ resonances, which include an angular-independent ($-1 \leftrightarrow 1$) transition that is in X band expected around 0.6 T [47]. Therefore, we attribute all the observed ESR components to the Mn$^{2+}$ and Mn$^{4+}$ ions. Moreover, we note that our calibration of the ESR intensity with a standard sample revealed a nice correspondence between the ESR susceptibility $\chi_{\text{ESR}}$ and the bulk susceptibility $\chi_b$, measured with a SQUID magnetometer (see Table II). This agreement proves that ESR detects all the spins in the Mn-doped SrTiO$_3$ samples and that all three distinct components of the ESR spectrum are intrinsic. Moreover, we find a relatively good correspondence (except for the A3N$_2$ sample) between the relative intensity $I_1$ and the occupancy of the A sites with Mn$^{2+}$ ions (see Table I). Therefore, the broad ESR component is mainly attributed to the Mn$^{4+}$ spins on the B site.

The temperature evolution of the three ESR components is shown in Fig. 4 for the B3O$_2$ sample, in which the broad C3 component is the most intense. The C1 sextet disappears below $\sim$120 K, similarly as observed before [27]. This can be attributed to freezing of the Mn$^{2+}$ ionic motion between the six energetically equivalent off-center positions within the A-site dedecahedron, which results in a much larger magnetic anisotropy and, consequently, in a more complicated anisotropy-split ESR powder spectrum below $\sim$120 K [27]. A gradual slowing down of the Mn$^{2+}$ hopping with decreasing temperature reflects already above 120 K in the increasing temperature reflects already above 120 K in the increasing

![Image of ESR spectra](image_url)

**FIG. 3.** (Color online) The room-temperature ESR spectra of 3% Mn-doped SrTiO$_3$ samples (circles) and their fits with a three-component model (see text for details). The three individual components are displaced vertically for clarity.

**FIG. 4.** (Color online) Temperature-dependent ESR spectra of the B3O$_2$ sample. The individual spectra are displaced for clarity. The lowest-field peaks of both sextets are marked by dashed lines. (b) The temperature dependence of the ESR linewidths of the three components. (c) Comparison of the inverse ESR susceptibility of the broad component $1/\chi_3 = 1/(I_1 \chi_{\text{ESR}})$ and bulk susceptibility $1/\chi_b$, measured in the field of 10 mT. Solid lines are linear fits of the high-temperature data.

**TABLE II.** Parameters of the three ESR components of the Mn-doped SrTiO$_3$ samples, their room-temperature ESR intensities $\chi_{\text{ESR}}$ and bulk susceptibilities $\chi_b$ measured by SQUID in the applied field of 10 mT.

<table>
<thead>
<tr>
<th>Component 1</th>
<th>Component 2</th>
<th>Component 3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sample</strong></td>
<td>$I_1$ (%)</td>
<td>$g_1$ a$_1$</td>
</tr>
<tr>
<td>A3N$_2$</td>
<td>26(5)</td>
<td>2.001 8.8</td>
</tr>
<tr>
<td>A3O$_2$</td>
<td>27(5)</td>
<td>2.000 8.7</td>
</tr>
<tr>
<td>B3N$_2$</td>
<td>4(2)</td>
<td>2.002 8.9</td>
</tr>
<tr>
<td>B3O$_2$</td>
<td>3(2)</td>
<td>2.002 8.8</td>
</tr>
</tbody>
</table>
narrowing, as the height is proportional to the spectrum height with doping is predominantly due to line intensity increases only slightly (note that the increase of the Weiss temperature of θ
= −15(5) K. The linewidth of the C3 component, on the other hand, is more or less temperature independent down to around 50 K and then increases notably down to 5 K. This linewidth increase can be interpreted as a fingerprint of a slowing down of spin fluctuations due to developing spin correlations. Once again, in accordance with the μSR results, we observe no anomalies in any of the ESR components (and in any of the investigated samples) around 40 K, where the spin-glass state was suggested to set in for the A-site-doped samples [16,17,31]. Our measurements thus provide a local-probe verification that such a spin state is not intrinsic and that the Mn-doped SrTiO3 samples rather remain paramagnetic down to the lowest temperatures.

In order to determine the origin of the broad ESR component that is dominant in all samples (see I3 in Table II), we have performed a doping-dependent study for all four different sample types. In Fig. 5, the ESR spectra in the doping range c = 1%–3% are shown for the families with the least intense (Aco2) and the most intense (Bco2) broad component. In all cases, we observe that the C3 linewidth notably and systematically narrows with the increasing doping concentration, e.g., from 40(2) to 28(2) mT for the B1O2 and B3O2 samples, respectively. At the same time, its relative intensity increases only slightly (note that the increase of the spectrum height with doping is predominantly due to line narrowing, as the height is proportional to 1/ΔB^2), mostly at the expense of the C2 sextet, while the relative intensity of the C1 sextet remains unchanged within the experimental error bars.

![Diagram](https://via.placeholder.com/150)

**FIG. 5.** (Color online) Doping-dependent normalized ESR spectra of the SrTiO3 samples (solid lines) at room temperature. The spectra corresponding to different concentrations are displayed for clarity. The thick dashed lines show the broad-line (C3 component) contribution to each spectrum, while the thin vertical lines indicate the lowest-field peaks of both sextets.

**IV. DISCUSSION**

The intensity of the unstructured broad ESR component C3 dominates the ESR spectra in all investigated samples. Although such component with a Lorentzian line shape is regularly observed in Mn-doped SrTiO3, especially for B-site doping [25,33,34,50], its origin remains unclear. Its alleged absence in some cases, e.g., as in Ref. [51], should be taken with caution, given the fact that due to its broadly distributed spectral weight with respect to the much narrower individual lines within sextets, it can easily be overlooked. It was ascribed before to the presence of the segregated MnTiO3 phase [33]. Since this phase orders magnetically at TN = 65 K, causing severe broadening of the ESR line close to TN and its disappearance below TN [52], while in our samples the width of the broad ESR line changes gradually and smoothly down to 5 K, this can not be the case here. Next, in reduced SrTi1−xMnxO3 samples, the broad ESR component was correlated with a ferromagnetic response, that was either claimed intrinsic [34] or extrinsic [51]. Again, our samples show no ferromagnetic characteristics.

In order to clarify its origin and the corresponding relevance for the magnetism of the Mn-doped SrTiO3, we have critically assessed possible ESR broadening mechanisms. As the isotropic exchange interaction J commutes with the S operator and therefore yields a delta-function ESR response, only a finite magnetic anisotropy gives a finite ESR linewidth [48]. First, we consider the case of well-isolated magnetic species (J ∼ 0). The dipolar interaction between magnetic moments is always present, therefore, we have calculated its contribution to the ESR linewidth. In the slow modulation limit (kB J ≪ gμB ΔB, where kB, μB, and B0 are the Boltzman constant, the Bohr magneton, and the applied magnetic field, respectively) the second moment of the absorption line arising from the secular part of the dipolar interaction [48]

\[ M_2 = \left( \frac{μ_0}{4\pi} \right)^2 3S(S + 1)(gμ_B)^4 \frac{1}{4} \left\{ \sum_k \frac{(3\cos^2 \theta_{jk} - 1)^2}{r_{jk}^3} \right\}_j \]

(5)

yields the ESR linewidth ΔB^d ≈ M_2/gμB. Here, the sum runs over all the neighboring sites of a chosen site j at the distance r_{jk} that are occupied by spins S, θ_{jk} is the angle that the r_{jk} vector makes with the external magnetic field, and (...)_j denotes powder averaging. In the case of diluted magnetic lattices, one can sum over all the lattice sites and the dilute-limit linewidth is then given by ΔB^d(c) = cΔB^d because M_2 is an additive quantity for the 1/r^3-type interactions [43]. We find the linewidths ΔB^d_A = 205 mT and ΔB^d_B = 134 mT in the limit of either fully occupied A sites with Mn^{2+} ions or B sites with Mn^{4+} ions, respectively. In the scenario (i) of diluted dopants, these then yield ΔB^d_A(3%) = 6 mT and ΔB^d_B(3%) = 4 mT, in disagreement with the enhanced width of the C3 component. The scenario (ii) of enhanced dipolar interaction within Mn-rich clusters of SrTiO3 was suggested before to explain the broad ESR component [25]. We note though that for dense magnetic clusters the ΔB^d_A = 205 mT...
and $\Delta B_{\text{g}} = 134$ mT linewidths should be observed. These exceed the experimental ones by an order of magnitude. Moreover, the Gaussian distribution of internal fields would be expected [48], which is clearly incompatible with the experimental line shape. Alternatively, reduced linewidths could occur if the B sites within clusters were only partially occupied, e.g., 22% magnetic-moment density on B sites would yield the ESR linewidth of 30 mT. However, even partially occupied clusters can not explain the substantial experimentally observed decrease of $\Delta B_{\text{g}}$ with increasing doping concentration. The latter would decrease the distance between magnetic moments and thus increase the average dipolar interactions. In scenario (iii), the magnetic anisotropy responsible for the width of the C3 ESR component is induced by oxygen vacancies. A vacancy in the B-site oxygen octahedron induces a large uniaxial single-ion anisotropy that was estimated as $D = 0.80$ K for the Mn$^{2+}$-Vo center in reduced SrTiO$_3$ single crystals [46]. Since $k_B D / g \mu_B \Delta B_{\text{g}} \sim 20$, isolated Mn$^{2+}$-Vo centers would yield a complex and much broader ESR spectrum.

The C3 ESR component thus rather speaks in favor of the scenario (iv) where strongly interacting magnetic moments on the B site are responsible for the observed behavior. In this case, large isotropic exchange ($k_B J \gg g \mu_B \Delta B$) leads to exchange narrowing and to a featureless Lorentzian line shape with the linewidth $\Delta B \approx M_2 / k_B J + g \mu_B J$ [48]. Considering the dipolar interaction within dense clusters (taking into account also the nonsecular terms that renormalize [48] $M_2$ by 10/3) we estimate the required exchange interaction as $J \sim 3$ K, while for the single-ion anisotropy scenario the required exchange coupling is of the order $J \sim k_B D^2 / g \mu_B \Delta B_{\text{g}} = 15$ K. Indeed, the former prediction, yielding the mean-field Weiss temperature $|\theta| = nS(S + 1)J / 3 \sim 20$ K, where $n = 6$ denotes the number of nearest neighbors in a cubic lattice, is in excellent agreement with the experimental value $\theta = -15(5)$ K.

In principle, the sizeable magnetic exchange ($J \sim 3$ K) that is needed to account for the broad ESR component could arise from two distinct origins: either from magnetic clusters, where sizable superexchange interaction between nearest-neighbor-occupied B sites was theoretically predicted [33] or from a donor-impurity mediated interaction [35] in the limit of diluted and homogeneously dispersed dopants. In order to further inspect the latter scenario, which stems from oxygen vacancies and has been recently claimed responsible for ferromagnetism of reduced Mn-doped SrTiO$_3$ crystals [34], we have performed DRS measurements, which can provide information on in-gap states for band-gap insulators such as SrTiO$_3$, with the band gap of 3.2 eV [53]. In Fig. 6, we show the DRS spectra $\{F(R)E\}^{1/2}$ as a function of the incoming-light energy $E$, which are obtained from the reflectance $(R)$ spectra of the samples by the Kubena-Munk transformation $F(R) = (1 - R^2)^{1/2}$ [54]. Similarly as in Ref. [34], we observed a pronounced dopant-induced in-gap state at around 2 meV that is absent in the undoped SrTiO$_3$ sample [34]. This state can be interpreted as a deep vacancy level, which has been predicted recently to be occupied by a single electron due to a strong Coulomb repulsion [55]. As such, in contrast to shallower donor-electron states with much larger Bohr radius (typically between 5 and 10 Å) observed in some other band-gap insulating oxides [35], this in-gap state is well localized [55] and can not provide extended magnetic coupling in a low-doping limit. Moreover, the effective coupling mediated by the donor electrons is always ferromagnetic because it is quadratic in the dopant-to-donor-electron magnetic interaction. This contradicts our experimental observations that reveal antiferromagnetic interaction between Mn ions contributing to the broad C3 ESR component.

We are therefore inclined to the scenario of dopant aggregation, where the enhanced ESR linewidth of the C3 component can be explained by an interplay of the dipolar and superexchange interactions within dense Mn-rich cluster of SrTiO$_3$. With increasing the doping concentration, the fraction of the isolated Mn$^{4+}$ ions on the B sites decreases (see the C2 component in Fig. 5), as one would expect. Moreover, the decreasing linewidth with increasing doping suggests that the exchange-narrowing mechanism is becoming more effective. Although ESR is not best suited for quantifying the cluster sizes, like some other nanocharacterization tools [5,38], this experimental finding is a likely fingerprint of nanosized clusters. Namely, for small clusters, an average exchange coupling of a particular Mn$^{4+}$ ion to its neighbors, being a scalar sum, is more affected by nonmagnetic Ti$^{4+}$ neighbors than the average dipolar interaction, resulting from a vector sum. This effect would, however, die out in bigger clusters where the percentage of the moments on the cluster surface becomes small compared to the moments residing inside the cluster. Thus, the scenario of a bulk phase segregation can safely be dismissed. The situation rather resembles other
diluted magnetic semiconductors, where nano-organization of dopants within the host matrix has been detected [5,38].

The apparent systematic mismatch between the relative ESR intensity of the C1 component \( I_1 \) (see Table I) and the site occupancy determined by x-ray absorption near-edge structure (XANES) spectroscopy [30] (see Table I) indicates that also Mn\( ^{2+} \) ions at the A sites aggregate, although only partially. Since the Mn[Mn]O\(_3\) cubic perovskite structure is unstable and a perovskite-type structure stabilizes only at high pressures and high temperatures [56], it would be structurally highly unfavorable if both Mn\( ^{2+} \) and Mn\( ^{4+} \) ions entered the same clusters. Therefore, a formation of separate MnTiO\(_3\) and SrMnO\(_3\) nanoclusters is expected. We note that the former compound in bulk adopts the ilmenite crystal structure [52] that is significantly different from the cubic perovskite structure of SrTiO\(_3\) as the Mn\( ^{2+} \) ions reside in oxygen octahedra. SrMnO\(_3\), on the other hand, can form a cubic perovskite polymorph [57]. This then probably explains the different tendency of Mn\( ^{2+} \) and Mn\( ^{4+} \) to aggregate. Quite importantly, we point out that the percentage of the aggregated Mn\( ^{2+} \) ions in SrTiO\(_3\) is correlated to the sample quality. There is a considerably larger mismatch between \( I_1 \) and the A-site occupancy [30], i.e., higher level of Mn\( ^{2+} \) aggregation in the A3N\(_2\) than in the A3O\(_2\) sample, while, on the other hand, the presence of structural defects determined by transmission electron microscopy (TEM) is much lower in A3N\(_2\), as its microstructure is more uniform [30].

Last, we return to the \( \mu \)SR results. Again, the homogeneous picture of the donor-electron-mediated exchange can not explain the rather slow fluctuations rates of local magnetic fields at the muon stopping sites (see Table I). The derived fluctuation rate \( \nu = 20 \text{ MHz} \) is orders of magnitude below the expected rate for exchange-coupled moments \( k_B J / h \sim 60 \text{ GHz} (J = 3 \text{ K}) \). In the inhomogeneous picture of clusters, on the other hand, the volume fraction of such clusters will be very small for the low doping concentrations of our samples. Therefore, the muons will predominantly couple to the diluted isolated moments outside the clusters, which fluctuate due to much smaller dipolar interactions. A small fraction of muons that do stop inside or close to clusters is likely responsible for the overestimation of the predicted LF decoupling at intermediate fields [Fig. 1(c)].

V. CONCLUSIONS

In conclusion, our muon spin relaxation and electron spin resonance investigations of Mn-doped SrTiO\(_3\) have demonstrated unambiguously that this system, independent of the nominal doping site and post-treatment atmosphere, intrinsically remains paramagnetic down to low temperatures. Both \( \mu \)SR depolarization curves as well as ESR spectra failed to detect any irregularities in the A-site-doped samples at the presumed spin-glass transition temperature around 40 K. In addition to several recent reports suggesting an extrinsic nature of ferromagnetism in many DMOs, our study, disproving the intrinsic spin freezing in the Mn-doped SrTiO\(_3\) materials, serves as a warning that the phase-segregation processes in not optimally processed materials can lead to very faint compositional inhomogeneities that are far below the limits of conventional analytical techniques. Moreover, we have discovered site-dependent aggregation of the dopants in Mn-rich clusters within the SrTiO\(_3\) matrix. Quite importantly, such spinodal decomposition is correlated to the quality of the sample’s microstructure. Moreover, it is less favorable for the Mn\( ^{2+} \) ions that partially remain isolated in diluted regions, as revealed by the remaining ESR sextet. Finally, our study has also unraveled the origin of the dominant broad ESR component commonly observed in the ESR studies of the Mn-doped SrTiO\(_3\) powders and crystals. This component emerges from the Mn-rich clusters of SrTiO\(_3\) and is due to exchange coupling and dipolar interactions between dopants.

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