Local spin susceptibility of the $S = \frac{1}{2}$ kagome lattice in ZnCu₃(OD)₆Cl₂

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(Received 12 March 2011; revised manuscript received 8 June 2011; published 21 July 2011)

We report a single-crystal ²D NMR investigation of the nearly ideal spin S = 1/2 kagome lattice $ZnCu_3(OD)_6Cl_2$. We successfully identify ²D NMR signals originating from the nearest neighbors of Cu^{2+} defects occupying Zn sites. From the ²D Knight-shift measurements, we demonstrate that weakly interacting Cu^{2+} spins at these defects cause the large Curie-Weiss enhancement toward T = 0 commonly observed in the bulk susceptibility data. We estimate the intrinsic spin susceptibility of the kagome planes by subtracting defect contributions, and explore several scenarios.

DOI: 10.1103/PhysRevB.84.020411

PACS number(s): 75.10.Kt, 76.60.-k

Persistent search for frustrated spin-liquid systems without a magnetically ordered ground state^{1,2} has recently resulted in the discovery of several model materials with spin S = 1/2, including the kagome,^{3–5} triangular,⁶ and fcc^{7,8} lattice systems. Intense experimental and theoretical efforts are underway to understand their ground-state properties, and to identify potential signatures of exotic spin-liquid states. Among the most promising candidates of spin liquids is herbertsmithite ZnCu₃(OH)₆Cl₂, which contains planes of a kagome lattice formed by Cu²⁺ ions with S = 1/2 (see Fig. 1 for the crystal structure).^{3,4,9–14} The antiferromagnetic Cu-Cu exchange interaction J is estimated to be as large as $J/k_B \sim 190$ K,⁴ yet there is a firm consensus that no hint of magnetic long-range order exists at least down to 50 mK.^{4,9}

Nonetheless, the mechanism behind the realization of such a nonmagnetic ground state in ZnCu₃(OH)₆Cl₂ has been controversial (see Ref. 15 for a recent review). The root cause of the controversy stems from the difficulty in pinpointing the location of defects, and understanding their potential influence on various physical properties. For example, magnetic Cu²⁺ defects occupying the nonmagnetic Zn^{2+} sites outside the kagome plane might induce a large Curie-Weiss contribution χ_{CW} to the bulk-averaged spin susceptibility χ_{bulk} , and mask its intrinsic behavior. In fact, χ_{bulk} always exhibits a large enhancement below about 50 K, as shown in Fig. 2(a). The observed behavior is different from the predicted behavior of χ_{kagome} for ideal kagome antiferromagnets with vanishing spin susceptibility at $T = 0.^{16-22}$ Furthermore, if nonmagnetic Zn^{2+} defects occupy the magnetic Cu^{2+} sites within the kagome plane, spin singlets may be induced in their vicinity toward T = 0¹² and help realize the nonmagnetic ground state. These uncertainties over the nature of defects prevented extraction of the intrinsic χ_{kagome} from χ_{bulk} . It is therefore necessary to untangle the effects of defects based on real-space probes.

In this Rapid Communication, we take advantage of recent successful growth of high quality single crystals of deuterized $ZnCu_3(OD)_6Cl_2$,¹⁴ and report a single-crystal NMR investigation. Thanks to very sharp NMR lines in a 45-mg single crystal, we identified ²D NMR signals originating from the 6 nearest-neighbor (NN) and 12 next-nearest-neighbor (NNN) ²D sites of Cu²⁺ defects occupying the Zn sites. From

their relative intensities, we show that magnetic Cu²⁺ defects replace nonmagnetic Zn²⁺ with a probability of $14 \pm 2\%$. A Curie-Weiss behavior of the NMR Knight shift ²*K* observed at the NN sites indicates that the large Curie-Weiss enhancement commonly observed for χ_{bulk} below ~50K arises from these defects. By subtracting the contribution of the Curie-Weiss contribution χ_{CW} from χ_{bulk} , we explore the intrinsic behavior of spin susceptibility of the kagome planes, χ_{kagome} .

In Fig. 3, we present representative ²D NMR line shapes observed in an external magnetic field of $B_{\text{ext}} = 8.4 \text{ T}$ applied along the c axis, i.e., normal to the kagome plane. The ^{2}D nucleus has a nuclear spin I = 1, and hence we would expect to observe a pair of NMR transitions between the nuclear spin state $I_z = \pm 1$ and $I_z = 0$, if there was no defect. In general, one can express the resonant frequency of each ²D site as $f_{\pm} = \gamma_n B_{\text{total}} \pm v_Q^c$, where the nuclear gyromagnetic ratio is $\gamma_n/2\pi = 6.53566$ MHz/T, and B_{total} represents the total magnetic field seen by each nuclear spin. The second term, $\pm v_{O}^{c}$, represents contributions of the c axis component of the nuclear quadrupole interaction for the aforementioned $I_z = \pm 1$ to $I_z = 0$ transitions. The ν_Q^c is proportional to the second derivative of the total Coulomb potential at the ²D sites, and small differences in the local structure lead to distinct values of the NMR line splitting $2\nu_O^c$.

In reality, we observe three pairs of ²D NMR signals, A_{\pm} , B_{\pm} , and C_{\pm} . This means that there are three distinct ²D sites in ZnCu₃(OD)₆Cl₂ with locally different environment. Based on the relative intensity of well-resolved peaks A_+ , B_+ , and C_+ at 295 K, we estimate the population of A, B, and C sites as $14 \pm 2\%$, $28 \pm 4\%$, and $58 \pm 4\%$, respectively. From the line splitting, we obtain $v_Q^c = (f_+ - f_-)/2 = 48$, 62, and 45 kHz for A_{\pm} , B_{\pm} , and C_{\pm} peaks, respectively. Within experimental uncertainties, we did not observe any noticeable temperature dependence in v_Q^c . Thus the temperature dependence of f_{\pm} arises from magnetic effects through $\gamma_n B_{\text{total}}$.

The B_{total} at each ²D site is a summation of the external field $B_{\text{ext}} = 8.4$ T and the hyperfine magnetic field B_{hf} from nearby electron spins polarized by B_{ext} , i.e., $B_{\text{total}} = B_{\text{ext}} + B_{hf}$. Since B_{hf} is proportional to *local* spin susceptibility χ_{spin} in the vicinity of the observed ²D site, one can gain insight into the position dependent χ_{spin} within the kagome plane by keeping track with the temperature dependence of f_{\pm} at different sites.

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FIG. 1. (Color online) Left: local configuration of ions surrounding the Zn^{2+} site in $ZnCu_3(OD)_6Cl_2$. Right: the *c*-axis view of the kagome plane.

A striking feature of Fig. 3 is that A_{\pm} peaks progressively shift to lower frequencies, and split off below 10 K. As shown below, this is because A sites are under the direct influence of the Cu²⁺ defect spins at NN Zn sites.

In Fig. 4, we summarize the temperature dependence of the ²D NMR Knight shift ² $K = B_{hf}/B_{ext} = (B_{total} - B_{ext})/B_{ext}$, which is related to local spin susceptibility as ² $K = \sum_i A_{hf}^{(i)} \chi_{spin}^{(i)} + K_{chem}$. $A_{hf}^{(i)}$ is the hyperfine coupling constant between the observed ²D nuclear spin and the electron spin at the *i*th Cu site in their vicinity, and $\chi_{spin}^{(i)}$ is the local spin susceptibility at the *i*th Cu site. $A_{hf}^{(i)}$ is negative in the present case, hence we reverse the vertical axis of Fig. 4. K_{chem} is a small temperature-independent chemical shift originating from the orbital motion of electrons. The results in Fig. 4 indicate that the Knight shift ²K(A) at the A sites is anomalously large compared with ²K(B) and ²K(C) at B and C sites. Furthermore, the temperature dependence of ²K(A)obeys a single Curie-Weiss law in the entire temperature range with a Weiss temperature, $\theta_A = -3.5$ K.

In Fig. 2, we compare the temperature dependences of ${}^{2}K(A)$ and χ_{bulk} . It is interesting to note that ${}^{2}K(A)$ asymptotes to χ_{bulk} at the low-temperature limit. In fact, the low-temperature behavior of χ_{bulk} can be approximated by a Curie-Weiss law $\chi_{CW} \sim 0.12/(T + \theta_{bulk})$ emu/mol f.u. with a vanishingly small Weiss temperature, $\theta_{bulk} = -1.2$ K; the extrapolation of χ_{CW} to higher temperature reproduces the temperature dependence of ${}^{2}K(A)$, as shown by a solid curve. The slight overestimation of $\theta_{A} = -3.5$ K in $B_{ext} = 8.4$ T by NMR over $\theta_{bulk} = -1.2$ K should be attributed to a saturation of defect spin polarization in high fields below $T \sim \mu_{B}B_{ext}/k_{B} \sim 5.6$ K. We confirmed that ${}^{2}K(A)$ at 4.2 K measured in $B_{ext} = 5.3$ and 3.2 T is indeed somewhat larger, and the field dependence is consistent with that of bulk magnetization.

Thus we have identified the source of the large enhancement of χ_{bulk} observed below 50 K as arising from defect moments neighboring ²D A sites whose behavior can be modeled with a Curie-Weiss law. The vanishingly small Weiss temperature implies that interactions among these defect moments are very weak. This weak coupling may explain the effectiveness of a



FIG. 2. (Color online) (a) (+): The bulk averaged magnetic susceptibility χ_{bulk} measured by superconducting quantum interference device (SQUID) in 0.2 T for a collection of small single crystals with random orientations. The solid curve represents extrapolation of the Curie-Weiss contribution χ_{CW} with $\theta_{\text{bulk}} = -1.2$ K estimated by fitting χ_{bulk} below 3 K. (•): ²D NMR Knight shift ²*K*(*A*) measured at the *A* site, which exhibits nearly identical temperature dependence as χ_{CW} . (o): The intrinsic spin susceptibility within the kagome plane, $\chi_{\text{kagome}} = \chi_{\text{bulk}} - \chi_{CW}$. (b) (o): the same χ_{kagome} plotted in (a). The solid curve through the data points represents the best fit with a small gap (see main text). The dotted curve shows the theoretical prediction (Ref. 18) with $J/k_B = 180$ K. (•): $\chi_{\text{kagome}}^{DF}$ estimated from the best fit of χ_{bulk} below 8 K to a summation of the *T*-linear term from Dirac Fermions and a Curie-Weiss term. (Δ): An alternative scenario of χ_{kagome}^c with a finite gapless excitation *c* at T = 0.

simple Curie-Weiss law, even though a distribution of coupling strengths would be expected for a random arrangement of defects. The next question to address then is where are these defects located? We recall that A sites account for $14 \pm 2\%$ of the overall ²D NMR signals, and the intensity of B sites is precisely twice greater, $28 \pm 4\%$. In view of the fact that each Cu^{2+} defect occupying the Zn sites has 6 NN and 12 NNN ²D sites (see Fig. 1), we assign A and B sites as the NN and NNN sites of the Cu^{2+} defects, respectively. These site assignments also imply that $14 \pm 2\%$ of the Zn sites in our single crystal are occupied by weakly interacting Cu^{2+} spins, and there is no ²D NMR signature of Zn²⁺ antisite defects occupying the Cu^{2+} kagome sites. Our findings are consistent with the recent sitespecific x-ray anomalous dispersion measurements combined



FIG. 3. (Color online) Representative ²D NMR line shapes observed in $B_{\text{ext}} = 8.4$ T applied along the *c* axis. We observe three pairs of quandrupole-split NMR lines, A_{\pm} , B_{\pm} , and C_{\pm} . Downward arrows mark the resonant frequencies f_{\pm} for A_{\pm} . Notice that the horizontal scale is changed for (b).

with Rietvelt refinement, from which the actual composition of our sample was estimated as $(Zn_{0.85}Cu_{0.15})Cu_3(OD)_6Cl_2$.²³

We gain additional insight into the nature of defects from the sign of the hyperfine interaction, A_{hf} . Our observation, ${}^{2}K(A) < 0$, implies that the hyperfine field B_{hf} at ${}^{2}D A$ sites, induced by the defect moments polarized along the c axis in B_{ext} , is negative. From the comparison of the temperature dependence of ${}^{2}K(A)$ and χ_{CW} , we estimate $A_{hf} \propto d({}^{2}K)/d(\chi_{CW}) = -24 \text{ kOe}/\mu_{B}$. On the other hand, in the present geometry employed for our NMR measurements, it is easy to show that the dipole hyperfine field from Zn sites, $A_{hf}^{\text{dipole}} = +0.85 \text{ kOe}/\mu_B$, has a *positive* component along the c axis. Fermi's contact hyperfine interaction induced by direct overlap of the ²D 1s orbital and Cu $3d^9$ orbitals at Zn sites is also positive. These considerations suggest that a sizable spin polarization obeying the same Curie-Weiss behavior also exists in the $Cu_3(OD)_3$ triangles within the kagome planes above and below the Zn defect sites; the negative hyperfine field at ²D A sites originates from these spin polarizations. This conclusion also explains the fact that $14 \pm 2\%$ of isolated Cu²⁺ defects alone would underestimate the magnitude of χ_{CW} by a factor of ~ 2 .

In contrast with the Curie-Weiss behavior of A sites, the Knight shift ${}^{2}K(C)$ at C sites is very small, and does not show



FIG. 4. NMR Knight shift ${}^{2}K$ at three distinct ${}^{2}D$ sites. Solid curve represents the best Curie-Weiss fit.

a large enhancement toward T = 0, as shown in Fig. 4. In other words, at least 58% of Cu sites within the kagome planes show no hint of enhancement of local spin susceptibility χ_{spin} toward T = 0. This means that the defect moments do not significantly couple to the Cu ions in the kagome plane beyond the NNN Cu sites, with which the 2D C sites are bonded. Since we cannot resolve the ²D B sites from C sites below ~50 K, 28% of Cu sites adjacent to B sites do not have a large enhancement of χ_{spin} , either, although we cannot entirely rule out a mild enhancement.

We have established that the Curie-Weiss contribution χ_{CW} to χ_{bulk} originates from the immediate vicinity of the Cu²⁺ defect moments occupying the Zn sites. This finding paves an avenue to analyze the intrinsic spin susceptibility within the kagome plane χ_{kagome} by subtracting χ_{CW} from χ_{bulk} , i.e., $\chi_{\text{kagome}} = \chi_{\text{bulk}} - \chi_{CW}$. In earlier studies of various quantum antiferromagnets, χ_{CW} was generally subtracted from χ_{bulk} to reveal intrinsic behavior, e.g., Ref. 24. Because of the uncertainties in the origin of defects, such analysis of χ_{kagome} has not been fully explored in the present case. In Fig. 2(a), we present $\chi_{\text{kagome}} = \chi_{\text{bulk}} - \chi_{CW}$, together with χ_{bulk} and χ_{CW} . We also present the same χ_{kagome} in Fig. 2(b) in an expanded scale.

Once we subtract χ_{CW} , a qualitatively different behavior of χ_{kagome} emerges; χ_{kagome} saturates below ~100 K, then decreases with temperature toward T = 0. The temperature dependence of χ_{kagome} is consistent with the theoretical prediction based on a high-temperature series expansion with $J/k_B \sim 180$ K as shown by a dotted curve. Our finding near T = 0 is also consistent with earlier reports based on ³⁵Cl and ¹⁷O powder NMR that χ_{spin} at some fraction of Cu sites decreases toward $T = 0.^{11,12}$ However, dramatic NMR line broadening hampered the efforts to probe the bulk-averaged behavior below 50 K in these earlier powder NMR studies.

The solid curve in Fig. 2(b) represents the best fit to an empirical formula $\chi_{\text{kagome}} = A \cdot \exp(-\Delta/k_B T)/(T + \theta_{\text{kagome}})$ with a small gap $\Delta/k_B \sim 7$ K and $\theta_{\text{kagome}} \sim 690$ K. The fit is reasonably good, but we should be cautious in interpreting its implications. Since we estimated χ_{CW} from χ_{bulk} below 3 K, and subtracted the former from the latter, in essence, we are forcing χ_{kagome} to reach zero at 3 K. In other words, the estimated values of χ_{kagome} set the lower bound of the intrinsic spin contributions at kagome sites. Thus we cannot rule out the possibility that this small gap is an artifact of our analysis. In this sense, we should consider $\Delta \sim 0.04J$ as the *upper bound* of any possible spin gap.

In fact, we noticed that the temperature dependence of χ_{kagome} near T = 0 is also consistent with a *T*-linear behavior predicted for gapless excitations of the Dirac Fermion in algebraic spin liquids, $\chi_{kagome} = A_{DF}T$, where $A_{DF} = 9.6N_A\mu_B^2k_B/J^2$ emu/K mol f.u.^{20,21} It is worth noting that the same Dirac Fermion model successfully accounts for the $C_{spin} \sim T^2$ behavior of the specific-heat data below 30 K with $J/k_B \sim 190$ K.^{20,21} Inspired by these promising signatures, we also fitted χ_{bulk} as a summation of the Dirac Fermion contributions $\chi_{kagome}^{DF} = A_{DF}T$ and defect contributions χ_{CW}^{DF} below 8 K, and estimate $\chi_{kagome}^{DF} = \chi_{bulk} - \chi_{CW}^{DF}$. It turned out that χ_{CW}^{DF} is nearly identical with the previous estimate, hence χ_{kagome}^{DF} shows very similar behavior, as shown in Fig. 2(b). The best-fitted value, $A_{DF} = 2.0 \times 10^{-4}$ emu/K mol f.u., implies $J/k_B \sim 220$ K. This is in reasonable agreement with $J/k_B = 180-190$ K mentioned above.

While the predictions based on Dirac Fermions with zero density of states at the Fermi energy account for both χ_{kagome} and C_{spin} fairly well, we caution that the nature of low-lying states of spin liquids is still controversial. For example, spin liquids with a spinon Fermi surface on triangular lattices are known to exhibit a finite density of states.^{6,25}

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An analogous scenario, however, seems highly unlikely in the present case, because C_{spin} exhibits positive curvatures in a broad temperature range below 30 K,⁴ instead of a *T*-linear behavior expected for a finite density of states. We estimate the upper bound of the *T*-linear term as $C_{\text{spin}} \sim 0.1 T$ J/mol f.u. K² below 1 K in 14 T.⁴ This translates to the upper bound of the constant term in the T = 0 limit of χ_{kagome} as $c = 1.4 \times 10^{-3}$ emu/mol f.u. We fitted the SQUID data to $\chi_{\text{bulk}} = c + \chi_{CW}^c$ below 3 K, and subtracted χ_{CW}^c from χ_{bulk} to estimate χ_{kagome}^c in the entire temperature range. The obtained results in Fig. 2(b) are similar to the previous two cases, except for the finite intercept at T = 0.

To summarize, we identified the source of a large enhancement of spin susceptibility in ZnCu₃(OH)₆Cl₂ as weakly interacting defect moments induced by Cu²⁺ in Zn sites and their immediate neighbors, and paved an avenue to analyze the bulk spin susceptibility data. By subtracting the defect contributions, we estimated an upper and lower bound of the intrinsic spin susceptibility within the kagome planes, χ^{c}_{kagome} (with a finite intercept) and χ_{kagome} (with a small gap), respectively. The reality must be somewhere in between.

T.I. thanks E. Sorensen, S.-S. Lee, P.A. Lee, Y. Ran, D. Freedman, and H. Eisaki for helpful communications, and acknowledges the financial support by NSERC and CIFAR. The work at MIT was supported by the US Department of Energy under Grant No. DE-FG02-07ER46134.

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