Quantum Tunneling and Relaxation in Mn$_{12}$ac Studied by Magnetic Spectroscopy

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(Received November 22, 2018)

In the last years molecular magnets comprising large clusters of coupled magnetic ions have attracted considerable interest as mesoscopic systems which exhibit new properties such as molecular magnetic bistability, macroscopic quantum tunneling of magnetization, quantum phase interference, etc. [3,4]. Among them clusters with a high-spin ground state and large anisotropy, such as Mn$_{12}$O$_{12}$(CH$_3$COO)$_{16}$(H$_2$O)$_4$2CH$_3$COOH-4H$_2$O (abbreviated as Mn$_{12}$ac) are of a paramount interest.

Mn$_{12}$ac cluster can be considered as nanoparticle with an effective magnetic moment of 20µ$_B$ (µ$_B$ is the Bohr magneton), corresponding to a collective spin $S = 10$ of the exchange coupled twelve Mn ions within the molecule. Due to the strong anisotropy, the magnetic moment freezes along one of the two easy directions at low temperatures. This is governed by a crystal field (CF) which splits the ground $S = 10$ multiplet and provides a quasi-doublet structure $|±m⟩$ of the lowest energy levels. The energy barrier $\Delta E \approx 60$ K between the two lowest lying states $|±10⟩$ determines the thermally activated relaxation of the magnetization in Mn$_{12}$ac [3,4]. Below a blocking temperature of about 3 K resonant quantum tunneling was discovered in the form of peaks in the relaxation rate and steps in the hysteresis loops at regular intervals of magnetic field corresponding to a coincidence of the CF energy levels [3,4]. Since the rate of the tunneling for a ground state is small in Mn$_{12}$ac the tunneling occurs via appropriate thermally excited CF states for temperatures above approximately 2 K resulting in a shortcut at the top of the barrier which accelerates the relaxation; at even lower temperatures pure quantum tunneling is dominant [3,4].

Various spectroscopic investigations have been applied to Mn$_{12}$ac, including electron-spin resonance (ESR) [5,6] and inelastic neutron scattering [7]. In particular, a new type of high-frequency ESR method based on a quasi-optical spectroscopy [8] made it possible to directly observe the CF transitions in an equilibrium state and to study in details their characteristics, including the line-shapes [3]. In this Letter we report on the high-frequency magnetic spectroscopy of non-equilibrium phenomena in Mn$_{12}$ac which allows to directly and independently determine the full set of magnetic characteristics.

Mn$_{12}$ac was pressed into a pellet with a thickness of 1.39 mm and a diameter of 10 mm. The high-frequency ESR experiments were performed by means of a quasi-optical coherent-source technique [10]. The transmission spectra Tr($ν$) of the pellet were measured with a linearly polarized radiation (its magnetic field denoted as $h$), in the frequency range $ν = 8$ to 12 cm$^{-1}$ at temperatures down to 1.8 K and in an external magnetic field $H$ up to 8 T applied perpendicular to the radiation $k$-vector (Voigt geometry). One frequency scan over the full range takes about 20 seconds.

Examples of the equilibrium Tr($ν$) spectra are shown in Fig. 1a. The absorption line at $ν_0 = 10$ cm$^{-1}$ corresponds to the $|±10⟩ → |±9⟩$ CF transitions at $H = 0$. It shifts to higher frequency $ν_H(H) ≈ ν_0 + γ|H_z|$ ($γ = gµ_B/h$), $H_z = H\cosθ_H$, $θ_H$ is the angle between $H$ and the easy $C_4$ z-axis, $g ≈ 2$) and changes its shape with increasing $H$, due to the inhomogeneous splitting of the degenerate $|±10⟩$ and $|±9⟩$ states in the polycrystalline Mn$_{12}$ac.

To study metastable states of Mn$_{12}$ac we have investigated the time dependence of the Tr($ν$) spectra. Applying an external field $H = +0.45$ T shifts the absorption line to the position $ν_H > ν_0$ (Fig. 1b); we then reverse the field (in about a minute), which leads to the line shift to the position $ν_L ≈ ν_0 - γ|H_z| < ν_0$. Starting from this point ($t = 0$) the intensity of the $ν_L$-mode decreases with time while that of the $ν_H$-mode recovers correspondingly.

The energy levels of Mn$_{12}$ac are determined by the effective Hamiltonian $\mathbf{H}$

$$H_{\text{eff}} = DS_z^2 + BS_4^4 + C(S_4^4 + S_4^4)/2 - gµ_B\mathbf{S} \cdot \mathbf{H}$$

(1)

where $\mathbf{S}$ is the effective spin of the ground $S = 10$ multiplet, and $D$, $B$, and $C$ are the CF parameters. At small fields the low energy levels are determined mainly by $H_z$: $E_m = Dm^2 + Bm^4 - gµ_BH_zm$, $m = 0, ±1, ±2, ±3$ (inset of Fig.1b). At low enough temperatures for $H_z > 0$ only
the $|+10\rangle$ quantum state is populated and we observe only one magnetic dipole transition $|+10\rangle \rightarrow |+9\rangle$. After reversing the field ($H_\perp < 0$) the state $|+10\rangle$ becomes metastable and its population relaxes to the new ground state $|-10\rangle$; the intensity of the transition $|+10\rangle \rightarrow |+9\rangle$ (at $\nu_L < \nu_0$) decreases while that for the transition $|-10\rangle \rightarrow |-9\rangle$ (at $\nu_H > \nu_0$) grows (Fig.1b). For our polycrystalline non-oriented sample a random orientation of the easy-axis results in the population of both $|+10\rangle$ and $|-10\rangle$ CF states and in an inhomogeneous broadening of the corresponding transitions by the applied field.

New features in the relaxation phenomena appear at lower temperatures, $T \lesssim 2$ K, as shown in Fig. 2. The relaxation occurs “inhomogeneously”; i.e., it has a faster rate near a particular frequency $\nu^{(1)}_t$ = 9.57 cm$^{-1}$ where a peak in the $\text{Tr(}\nu\text{)}$ spectrum appears accompanied by a minimum developing at $\nu^{(1)}_t$ = 10.47 cm$^{-1}$. This new phenomenon can be called magnetic hole burning in analogy to spectral hole burning in dielectrics [10]. We ascribe this fast relaxation near the $\nu^{(1)}_t$ to a resonant quantum tunneling in those crystallites whose orientation relative to $\textbf{H}$ satisfy the level crossing condition $h\nu^{(1)}_t(m) = h\nu_0 \pm g\mu_B H_{\text{cross}} = h\nu_0 \pm [-D - B(m^2 + (m - 1)^2)]$.

For a quantitative description of the observed relaxation the effective magnetic permeability is calculated using (a) the non-equilibrium populations of the low CF states $|\pm 10\rangle$ and their evolution (relaxation) with time and (b) the inhomogeneous broadening of the CF transitions in the magnetic field due to the random orientation of the crystallites. Firstly, we analyze a behavior for a single crystallite. Considering only two $|\pm 10\rangle \rightarrow |\pm 9\rangle$ CF transitions and introducing a non-equilibrium normalized populations $\rho_m(t)$ we generalize the ordinary equilibrium permeability [8] and obtain its transverse components:

$$\mu_{xx,yy}(\nu, t) \equiv \mu_\perp(\nu, t) = 1 + \Delta \mu^+(\nu, t) + \Delta \mu^-(\nu, t) \quad (2a)$$

$$\mu_{xy,yx}(\nu, t) \equiv \pm \mu_\parallel(\nu, t) \equiv \pm \sqrt{\Delta \mu^+(\nu, t) - \Delta \mu^-(\nu, t)} \quad (2b)$$

where

$$\Delta \mu^\pm(\nu, t) = \Delta \mu_0 \left[ \frac{\nu_0}{\nu_0(\rho_{\pm 10} - \rho_{\pm 9})} R_\pm(\nu) \right]. \quad (3)$$

are the contributions of the $|\pm 10\rangle \rightarrow |\pm 9\rangle$ CF transitions to the permeability with its static value at $T = 0$, $H = 0$ being $\Delta \mu_0$; a Lorentzian lineshape $R_\pm(\nu)$ is assumed for simplicity; $h\nu_\pm = E_{\pm 9} - E_{\pm 10} = h\nu_0 \pm g\mu_B H_\perp$. The equilibrium state is given by $\rho_m(t \to \infty) = \rho_m^\infty = \exp[-\beta E_m]/Z$ with $Z = \sum \exp[-\beta E_m]$; $\beta = 1/k_B T$, and $\sum \rho_m(t) = 1$.

In the thermally assisted regime, the relaxation of the populations $\rho_m(t)$ takes place via excited states and is determined by the complex master equation for the populations of all CF states [12]. In this case $\Delta \mu^\pm(\nu, t)$ can be reduced to the simple relaxation equation $\Delta \rho(t) = -(\Delta \rho - \Delta \rho^\infty)/\tau$ for $\Delta \rho = \rho_{+10} - \rho_{-10}$ where $\Delta \rho^\infty = \rho_{+10}^\infty - \rho_{-10}^\infty$ is the difference of the equilibrium population after the field is reversed. The solution $\Delta \rho(t) = \Delta \rho^\infty [1 - 2 \exp\{-t/\tau\}]$ for the initial condition $\Delta \rho(0) = \Delta \rho^0 = -\Delta \rho^\infty$ leads to the population differences

$$\Delta \rho_{\pm}(t) \equiv \rho_{\pm 10} - \rho_{\pm 9} = \Delta \rho^\infty + (\Delta \rho^0 - \Delta \rho^\infty) e^{-t/\tau} \quad (4)$$

which enter Eq. (2); here $\Delta \rho^{\infty}_{\pm} = \{\exp[\beta E_{\pm 10}] - \exp[-\beta E_{\pm 10}]/\exp[-\beta h\nu_\pm]\} [\exp[-\beta h\nu_\pm]/2 + \exp[\beta h\nu_\pm] + \exp[-\beta h\nu_\pm]]$, where all quantities refer to the reversed field. The derivation assumes thermal equilibrium between $|\pm 10\rangle$ and $|\pm 9\rangle$ states due to fast relaxation via phonons, implying $\rho_{\pm 9} = \rho_{\pm 10} \exp[\mp \beta h\nu_\pm]$. For low temperatures ($\beta h\nu_\pm >> 1$), $\rho_{\pm 9}$ is negligible and Eq. (4) simplifies to $\Delta \rho_{\pm}(t) \approx \rho_{\pm 10}(t) \approx \rho_{\pm 10}^\infty + (\rho_{\pm 10} - \rho_{\pm 9}^\infty) \exp[-t/\tau]$.

All previous consideration refers to a single crystal. For polycrystalline samples we have to take into account the inhomogeneous broadening of the lines due to different splittings of the doublets. By averaging the permeability [Eq. (3)] of the crystallites with respect to their orientations we obtain the effective permeability $\mu^{\text{eff}}(\nu, t)$ with the non-zero components $\mu^{\text{eff}}_{xx,yy}(\nu, t) = \langle \mu_{\perp}(\nu, t)(1 + \cos^2 \theta_H) \rangle/2$, $\mu^{\text{eff}}_{xy,yx}(\nu, t) = \mp i(\mu_{\parallel}(\nu, t) \cos \theta_H)$, and $\mu^{\text{eff}}_{zz}(\nu, t) = \langle \mu_{\perp}(\nu, t) \sin^2 \theta_H \rangle$ for the $\textbf{H}$ oriented along the $z$-axis. Using $\mu^{\text{eff}}(\nu, t)$ and the Fresnel formulas for the transmission coefficient [17] we fit the zero-field spectrum and obtain $\nu_0 = 10.02$ cm$^{-1}$, $\Gamma = 0.18$ cm$^{-1}$, and $\Delta \mu_0 = 0.0107$, in accordance with our previous results [8]. These parameters also describe the spectra in the magnetic field [18] in all details, including the dramatic changes of the lineshape (Fig. 3a).

The relaxation phenomena observed in our $\text{Tr(}\nu\text{)}$ spectra at various temperatures can be self-consistently described using the theory of the magnetization relaxation in Mn$_2$ac [12–16] based on phonon-assisted spin tunneling induced by forth order CF and transverse magnetic field. To determine the relaxation rate $\tau(T, H, \cos \theta_H)$ we have numerically diagonalized the master equation for the density matrix $\rho = \rho_0 + \rho_1$ in the space of the ground $S = 10$ multiplet, following the procedure suggested by Ref. [17]. In calculations we used the CF parameters of Eqs. (1) $D = -0.389$ cm$^{-1}$, $B = -7.65 \times 10^{-4}$ cm$^{-1}$, $|C| = 5.0 \times 10^{-5}$ cm$^{-1}$ from [18] and parameters of the spin-phonon interaction determined via the main CF term $DS^2$.

In Fig. 3 the angular dependence of the relaxation time is plotted for two different temperatures and fields used in our experiments (Figs. 1c and 3). The relaxation goes fastest for $\cos \theta_H = 0$ because the energy levels on the both sides of the barrier coincide: a situation, however, not observed in Voigt geometry. For fields exceeding the value of the first level crossing ($\sim 0.44$ T), additional resonance minima arise with a fine structure related to the slightly different resonance conditions for various exciting levels due to the fourth order CF contribution B.
As an example, Fig. 1b shows the relaxation observed in the Tr(ν) spectra at \( H = 0.45 \) T and \( T = 2.6 \) K which are well described by our model. We note that a weak resonance near \( \cos \theta_H = 0.97 \) due to avoided level crossing of the CF states \( m = 4, -3 \) (or \( m = 3, -4 \)) does not exhibit in the spectra due to the averaging in the polycrystalline sample. The relaxation rate strongly increases with magnetic field because the effective barrier decreases with both (longitudinal and transverse) components \( \delta \); there are also significant contributions from the tunneling processes which are enhanced because the tunneling splitting increases with the transverse field.

These resonance effects are best seen at low temperatures, when the thermal relaxation over a barrier is suppressed. Fig. 2 shows a set of resonances in the relaxation rate at \( T = 1.96 \) K which correspond to pairs of states with an avoided level crossing. Using this \( \tau(\cos \theta_H) \), the time evolution of the Tr(ν) spectra is calculated; as seen from Fig. 3 the agreement is excellent. The main contribution to the resonance relaxation comes from the thermally assisted tunneling via \((5,-4), (6,-5)\) and in less degree \((7,-6)\) states (or corresponding states with the opposite signs); they determine the frequency of the burned spectral hole. For these avoided level crossings the resonance lineshape is determined by a truncated Lorentzian \([1, 13]\) with the effective linewidth determined by the tunneling splitting (0.23, 0.015, and 5.0 × 10^{-4} cm^{-1}, respectively) which is much larger than the corresponding phonon linewidth (ranging from 5 × 10^{-6} to 3 × 10^{-4} cm^{-1}). The linewidth (0.12 − 0.18 cm^{-1}) of the zero-field-resonance absorption \( \nu_0 \) exceeds the corresponding phonon linewidths considerably. It is mainly determined by inhomogeneous broadening and thus serves as a measure of the disorder effects \([3]\), which could originate from CF and g-factor distributions (D-strain and g-strain) and random magnetic dipolar fields as was shown by ESR \([22]\).

The problem of shape of the tunneling resonance and the effect of inhomogeneous broadening on it are presently subject of considerable interest and discussions \([12, 14, 24]\), many issues, however, remain open. One of them could be related to the obvious discrepancy between the observed inhomogeneous character of the zero-field modes broadening \([20]\) and Lorentzian shape of the tunneling resonances observed at temperatures \( T \gtrsim 2K \) \([14, 21]\) and described theoretically in the thermally assisted regime \([4]\). One may expect that a large inhomogeneous broadening of the levels (i.e., larger than the tunneling splitting) reduces the tunneling amplitudes and smears of the resonances, resulting in suppression of the relaxation peaks and in change of their Lorentzian shape \([13]\). Suggesting that the main contribution to the line broadening comes from the CF dispersion and assuming its Gaussian distribution \([22]\), we can evaluate the dispersion \( \delta D = 0.004 \) cm^{-1} of the main quadratic term \( D \) in the Hamiltonian \([4] [23]\). This yields a smearing of the tunneling level differences \( \delta(E_m - E_{m+1}) = \delta D(2m-1), \) for instance, of 0.035 − 0.05 cm^{-1} for \((5,-4), (6,-5),\) and \((7,-6)\) states. From a comparison with the calculated tunneling splittings we can conclude that the inhomogeneous level broadening does not effect the main resonance \((5,-4),\) smears the \((6,-5),\) but suppresses the \((7,-6)\) and higher order resonances. A numerical simulation including an additional averaging with respect to the CF distribution confirms this conclusion. Thus, if the tunneling splitting is sufficiently large for states contributing to the thermally assisted tunneling, the lineshape of the corresponding resonance remains Lorentzian.

In summary, we utilized a novel optical method of high frequency magnetic spectroscopy to study the relaxation phenomena in Mn_{12}ac clusters, including the resonance quantum tunneling along with the CF transitions and their lineshapes and linewidths. At elevated temperatures only frequency-homogenous relaxation due to mainly thermal activation is observed, whereas for \( T \lesssim 2 K \) the thermal relaxation is significantly suppressed and resonant quantum tunneling dominates; in this range the effect of magnetic hole burning is observed. We have developed a microscopic model which allows to quantitatively describe all observed effects, taking into account the orientational distribution of the relaxation time and to determine the most effective relaxation channels and to estimate the effect of CF inhomogeneity on the resonant tunneling.

We thank the Prof. N. Karl and the Materials Lab for synthesizing the Mn_{12}ac. We acknowledge support from the Deutsche Forschungsgemeinschaft (DFG). This work was supported in part by RFBR (No. 02-02-16597).

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Reversing the field we observe a weak asymmetry of the spectra with respect to the zero-field line position \( \nu_0 \) which indicates a small internal dipolar field \( \lambda M \approx 0.04 - 0.05 \) T; we have included it in our calculations.

FIG. 1. Transmission spectra of polycrystalline 1.39 mm thick pellet of Mn\(_{12}\)ac measured in an external magnetic field \( \mathbf{H} \perp \mathbf{h} \perp \mathbf{k} \). (a) Shift of the absorption line with increasing magnetic field. The solid lines represent Lorentzian fits with the distribution of the crystallites orientation taken into account. (b) Time evolution of the absorption line after the magnetic field is reversed from +0.45 T to −0.45 T. The lines represent the transmission for the calculated angular distribution of the relaxation time \( \tau(\cos \theta_H) \) shown in Fig.3. The arrows indicate the positions of line at \( \mathbf{H}=0, \pm 0.45 \) T. Inset: the energy levels of a Mn\(_{12}\)ac single crystal for a magnetic field \( \mathbf{H} \parallel C_4 \); the arrows show the \( |\pm 10\rangle \rightarrow |\pm 9\rangle \) transitions before and after the field inversion.

FIG. 2. Time evolution of the transmission spectra at \( T = 1.96 \) K after the magnetic field (\( \mathbf{H} \perp \mathbf{h} \perp \mathbf{k} \)) is reversed from +0.9 T to −0.9 T. The narrow absorption dip within the broad feature is due to resonant quantum tunneling between pairs of coinciding energy levels. The lines correspond to the transmission spectra for calculated distribution of the relaxation time \( \tau(\cos \theta_H) \) shown in Fig.3. Also shown is the zero-field absorption line.

FIG. 3. Calculated distribution of the phonon-assisted spin-tunneling relaxation time \( \tau(\cos \theta_H) \) as a function of the angle between the magnetic field \( \mathbf{H} \) and the \( C_4 \) axis of the crystallites at \( T = 2.6 \) K and \( T = 1.96 \) K for certain values of the applied magnetic field. The numbers at the resonance minima indicate pairs of tunneling CF states.
\section*{Fig. 1}
$H = 0.9 \text{T}$

$\nu = \nu_0 (1)$

$t = 0$

$285 \text{ min}$

$75 \text{ min}$

$H = 0 \text{T}$

$T = 1.96 \text{ K}$

$\text{Mn}_{12} ac$

$\nu_{-}^{(1)}$ $\nu_0$ $\nu_{+}^{(1)}$

$\nu = \nu_0 (1)$

$75 \text{ min}$

$285 \text{ min}$

$t = \infty$

$H = 0.9 \text{T}$

Transmission coefficient

Frequency (cm$^{-1}$)

Fig. 2
Fig. 3

Relaxation time (s)

$\cos \theta_H$

Mn$_{12}$ac

$T = 1.96 \, \text{K}$
$H = 0.9 \, \text{T}$

$T = 2.6 \, \text{K}$

$H = 0.2 \, \text{T}$

$\cos \Theta_H$