

Molecular rotors in a metal-organic framework: muons on a hyper-fast carousel.

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Abstract

Using muon-spin spectroscopy, we study the exceptional dynamical properties of rotating molecular struts engineered within a Zn-based metal-organic framework at cryogenic temperatures, where the internal motions of almost any other organic substance are quenched. Muon-spin spectroscopy is particularly suited for this aim, as the experimental evidences suggest one implantation site for the muons on the stator and – at least – another implantation site directly onto the rotating moiety. The dynamics of the molecular rotors is characterized by the exceptionally low activation energy $E_A \sim 30 \text{ cal mol}^{-1}$. At the same time, we evidence a highly unusual dependence of the dipolar interaction of muons with nuclear magnetic moments on the rotors, suggesting a complex influence of the rotations on the muon implantation and diffusion.

Keywords

Muon-spin spectroscopy, Molecular rotors, Metal-organic frameworks.

Research on molecular machines, rotors and switches is a very active field lying at the boundary between chemistry, physics and materials science whose ultimate aim is to control the mechanical motion of nanoscopic objects in order to implement useful tasks at the nanoscale.¹⁻⁸ The realization of rotational movements for molecules along controlled axes is of significant interest for most applications in this respect.^{9,10} In particular, the design and synthesis of materials endowed with fast dynamics made the prospects for several applications – such as selective sensors, gas sorption and release-on-command to switchable ferroelectrics – much more concrete.^{11,12}

For this aim, the recent use of porous materials as a new platform has extended the perspectives of fast molecular rotors to the crystalline solid state.¹²⁻¹⁶ Within this context, researchers have devoted particular interest to metal-organic frameworks (MOFs), which are engineered by self-assembly of rotor-containing ligands and integrate both stator and rotary elements in the same crystal structure. The highly porous nature of MOFs retains the free volume needed for the rotary motion to occur – at the same time, the accurate design of the surrounding crystal architecture and the low torsional energy profile about pivotal bonds are necessary to push the motional behaviour to its limits. By means of these approaches, ultra-fast dynamics comparable to liquid matter was demonstrated recently at cryogenic temperatures with characteristic activation energies markedly smaller than the thermal energies at ambient conditions.¹⁷⁻¹⁹

The in-depth knowledge of dynamical processes in solids is challenging – especially in the presence of fast motional processes. Positive muons are very convenient tools for this aim, since they act as highly sensitive local probes of microscopic fluctuating magnetic fields.²⁰⁻²⁵ Thus, muon-spin spectroscopy (μ SR) has been a valuable tool to reveal molecular reorientation in liquid-crystalline materials and molecular dynamics in general.²⁶⁻³¹ In this work, we report on the first μ SR study of the unique dynamical properties of molecular

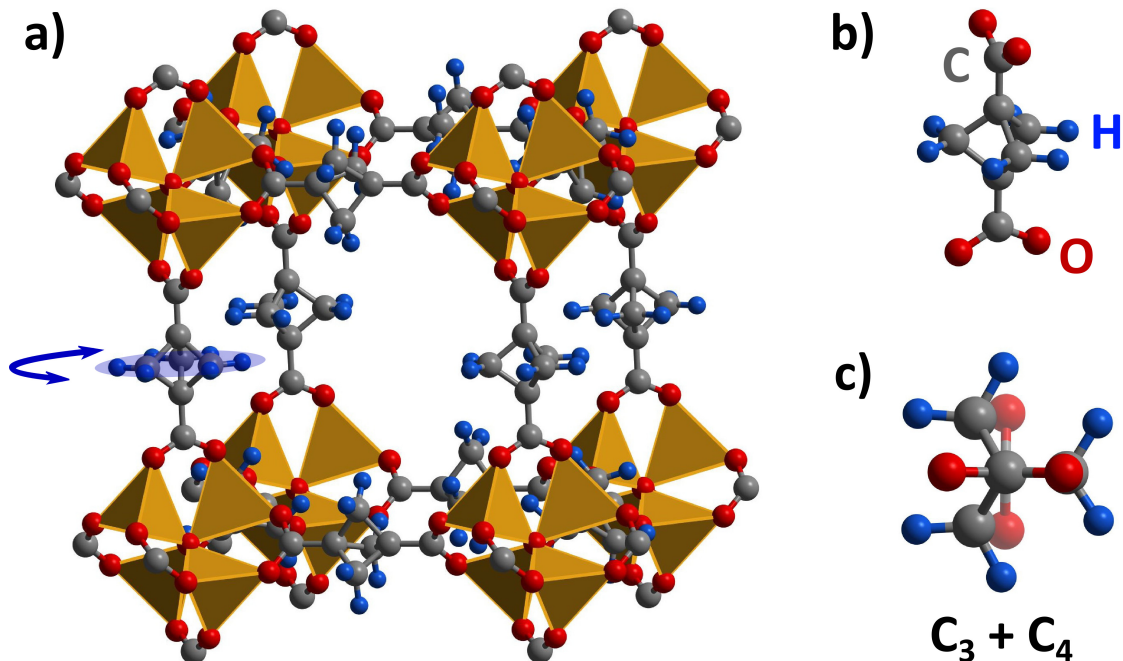


Figure 1: a) Cubic crystalline structure of the investigated MOF showing the rotating bicycle unit suspended within the Zn-based stators. The bicyclo-[1.1.1]-pentane-1,3-dicarboxylate moiety is also viewed b) parallel and c) perpendicular to the rotation axis.

rotors engineered in MOFs, focussing on a Zn-based MOF containing bicyclo-[1.1.1]-pentane-1,3-dicarboxylate struts.¹⁹ The cubic structure of this compound (Fig. 1a) contains nodes made by four Zn atoms with tetrahedral geometry coordinated with six bicyclopentane-dicarboxylic acid ligands. The nodes act as stator for the rotary motion of the bicyclo-[1.1.1]-pentane rotor. Geometrical frustration arises from the C_3 symmetry of the rotor conflicting with the crossed C_4 arrangement of the two neighbouring carboxylate groups (Figs. 1b and 1c). This delicate balance of different symmetries favours the generation of very shallow energy minima associated with the orientation of the bicyclo-[1.1.1]-pentane unit whose rotary motion – as a result – is characterized by a virtually complete compensation of the energy barrier. Remarkably, the dynamics is preserved down to cryogenic temperatures where almost any organic substance is solid and internal motions are quenched, exception being proton exchange and small moieties, such as amine and methyl groups. Details on the synthesis and characterization of the described MOF are reported in the supporting

information (Figs. SI1, SI2 and SI3).

Critical aspects of μ SR are the determination of the muon implantation crystallographic sites and the possible influence of the implanted muons on the properties of the investigated materials. For the measurements we are currently reporting on, we have evidences of at least two inequivalent implantation sites. One site is possibly located on the stator and muons implanted therein are sensitive to the rotational dynamics of the rotor as a whole, evidencing a value $E_A \sim 30 \text{ cal mol}^{-1}$ for the characteristic activation energy for the rotations – i.e., the rotors’ mobility. Although higher than the NMR estimate, likely because of the perturbation induced by the muons on the motions, this exceptionally low value is a benchmark for the easiness of rotation of the studied molecular rotors. On the other hand, the signal from those muons implanted in the other site(s) evidences well-defined coherent oscillations arising from the dipolar interaction with the nuclear magnetic moments of hydrogen atoms. This unambiguously locates the implantation site(s) directly onto the rotating moiety. The unprecedented evolution of the oscillating signal upon increasing temperature demonstrates that the implantation of muons onto the rotors is strongly affected by the onset of the rotary dynamics, which favours muons’ diffusion in turn.

As discussed in the supporting information, the main quantity of interest in a μ SR experiment is the so-called spatial asymmetry in the muons’ radioactive decays. It can be shown that this quantity is directly proportional to the spin autocorrelation function for the muon and, accordingly, a time-resolved measurement of the decay asymmetry is equivalent to measuring the time-evolution of the spin depolarization of the muons subject to the local static and dynamical magnetic fields probed in the investigated material. We report representative results for the muon depolarization under conditions of zero external magnetic field (ZF) in Fig. 2, evidencing a strong dependence of the experimental results on temperature. The value of the full initial asymmetry ($\sim 0.09 - 0.12$) is quite low compared to the maximum instrumental value (~ 0.25) regardless of the temperature value. We distinguish three different regimes. In the high-temperature range (HT, $T \gtrsim 13 \text{ K}$), the ZF

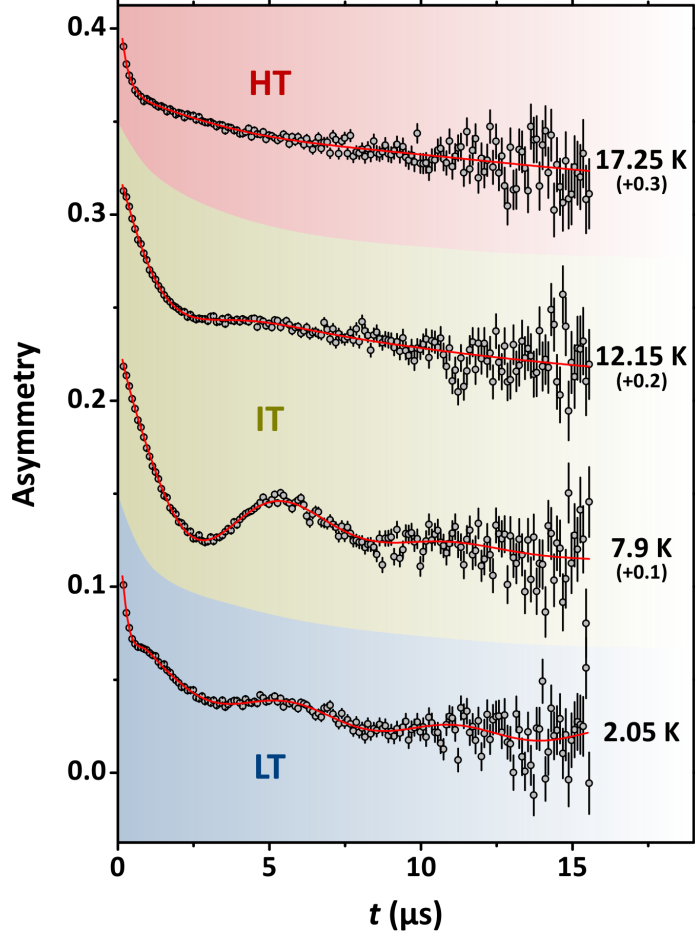


Figure 2: Representative results of ZF- μ SR at different temperature values in the low- (LT), intermediate- (IT) and high-temperature (HT) regions. The curves have been shifted vertically by the amounts in the parentheses for the aim of clarity. The continuous lines are best-fitting functions according to Eq. 1 for the data at 7.9 K, 12.15 K and 17.25 K. For the measurement at 2.05 K, the continuous line is a best-fitting function according to the H- μ model discussed in the supporting information (see Eq. SI11).

asymmetry function is reproducible as the sum of two decaying contributions with markedly different time constants ($\sim 6 \mu\text{s}^{-1}$ and $\sim 0.05 \mu\text{s}^{-1}$, respectively). Upon cooling the system, in the intermediate-temperature range (IT, $8 \lesssim T \lesssim 13$ K), the initial exponentially-decaying component evolves progressively in a well-defined coherent oscillation. Suddenly, below the temperature value $T \simeq 8$ K (low-temperature range, LT), the amplitude of the oscillating fraction gets unevenly redistributed between two cosine-like signals with different frequencies.

Based on the qualitative arguments above, we fitted the following function

$$A^{\text{ZF}}(t) = \sum_{i=1}^2 A_{f_i}^{\text{ZF}} \cos(\gamma_\mu B_i t) e^{-\lambda_{f_i}^{\text{ZF}} t} + A_s^{\text{ZF}} e^{-\lambda_s^{\text{ZF}} t} \quad (1)$$

to the experimental data in ZF conditions. Here, A^{ZF} is the total asymmetry of the muons' decays as a function of time t , split into different components with amplitudes $A_{f_i,s}^{\text{ZF}}$. In general, the two f components (labelled by the index i) are exponentially-damped cosine functions, where the angular frequency is written as the magnetic field at the muon site B_i times the muon gyromagnetic ratio $\gamma_\mu = 2\pi \times 13.554 \text{ rad ms}^{-1} \text{ G}^{-1}$ and where $\lambda_{f_i}^{\text{ZF}}$ are the associated damping rates. Finally, the long-tail relaxing component is damped exponentially with rate λ_s^{ZF} . In order to match the experimental findings described above, we fix $B_1 = 0$ in the HT region and $A_{f_2}^{\text{ZF}} = 0$ in both the IT and HT regimes. We obtain good fitting results (statistical $\chi^2 < 1.1$) throughout the whole temperature range.

In the following we will focus also on the effect of external magnetic fields H^{LF} parallel to the muons' spin at the moment of implantation – the so-called longitudinal field (LF) geometry. We report representative experimental curves in LF geometry in the insets to Fig. 3 and Fig. 4. Under the effect of a longitudinal field, we observe that the depolarization can be described as the sum of two exponentially-decaying functions with markedly different time constants – however, as thoroughly discussed later on in the text, only the long-tail component is observed in the LT regime. To reproduce these observations, we refer to the following expression

$$A^{\text{LF}}(t) = A_f^{\text{LF}} e^{-\lambda_f^{\text{LF}} t} + A_s^{\text{LF}} e^{-\lambda_s^{\text{LF}} t} \quad (2)$$

as best-fitting function for the depolarization curves at all temperatures, provided that $A_f^{\text{LF}} = 0$ within the LT region. The meaning of symbols in Eq. 2 can be derived straightforwardly based on the discussion of Eq. 1.

The central assumption in our analysis is that the different contributions in Eq. 1 and Eq. 2 are associated with signals from inequivalent implantation sites for the muons. We

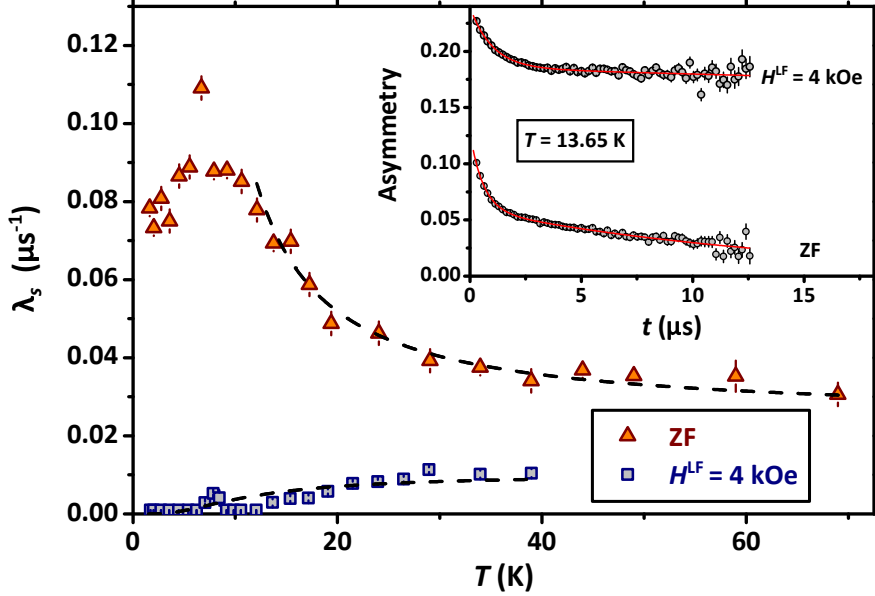


Figure 3: Main panel: temperature dependence of the relaxation rate λ_s under ZF conditions and for $H^{\text{LF}} = 4$ kOe. The dashed curves are the results of a simultaneous best-fitting procedure to both ZF and LF data based on Eqs. 3 and 4. Inset: representative depolarization curves at $T = 13.65$ K under conditions of ZF and LF ($H^{\text{LF}} = 4$ kOe). The continuous lines are best-fitting curves based on Eq. 1 and Eq. 2, respectively.

start from the discussion of the long-tail exponentially-damped component on temperature, which is preserved qualitatively unmodified across the whole investigated temperature range. Indeed, we find $A_s^{\text{ZF}} \simeq 0.05 - 0.06$ for the amplitude of this component at all the investigated temperatures. We report in Fig. 3 the temperature dependence of the associated relaxation rate λ_s^{ZF} as well as λ_s^{LF} for the longitudinal field $H^{\text{LF}} = 4$ kOe. The observed trend can be interpreted as being due to dynamical influences on the muon depolarization via spin-lattice-like relaxation processes. In the fast-fluctuations regime, the relation

$$\lambda_s(T) = 2\gamma_\mu^2 \Delta^2 \frac{\tau_c(T)}{1 + \omega_\mu^2 \tau_c^2(T)} \quad (3)$$

holds²³ where τ_c is the correlation time for the dynamics being probed by the muons, Δ is the root-mean-square amplitude of the fluctuations of the local magnetic fields and $\omega_\mu = \gamma_\mu H^{\text{LF}}$

is the Larmor frequency for the muon. Assuming that

$$\tau_c(T) = \tau_0 \exp(E_A/k_B T), \tag{4}$$

where $k_B = 1.38 \times 10^{-16}$ erg K⁻¹ is the Boltzmann constant, it is possible to quantify E_A – i.e., the characteristic activation energy for the rotations – and the correlation time at infinite temperature τ_0 by performing a simultaneous fit to the ZF and LF data for $\lambda_s(T)$ based on Eq. 3 and Eq. 4 for temperatures above $T \sim 10$ K. The results of the fitting procedure are shown as dashed lines in the main panel of Fig. 3. Based on our results, we estimate the remarkably low value $E_A = 30 \pm 2$ cal mol⁻¹, as well as $\tau_0 = 3.5 \pm 0.5$ ns. We argue that the dynamical process probed by the muons is the rotation of the bicyclo-[1.1.1]-pentane rotors and, accordingly, both E_A and τ_0 are characteristics of this rotary motion.

Some comments are in order at this stage. We report the recovery of the muon depolarization under the effect of an increasing longitudinal magnetic field in the supporting information (see Fig. SI4). Based on the observed recovery of the A_s amplitude as a function of the field intensity, we can associate this signal unambiguously to a muonium state – i.e., to the ~ 80 % of incoming muons forming a bound μ^+e^- state after thermalizing in the sample (see the supporting information).³² Additionally, we argue that these muons should be stably localized on the stators and probe via dipolar interactions the time-modulated magnetic fields generated by the moving rotors.

It is also meaningful to compare our results from μ SR with those collected by ¹H-NMR in the absence of implanted muons.¹⁹ The values for both E_A and τ_0 estimated by muons are higher than the estimates made by means of the NMR spin-lattice relaxation rate – in fact, $E_A^{\text{NMR}} \simeq 6$ cal mol⁻¹ and $\tau_0^{\text{NMR}} \simeq 4$ ps.¹⁹ In this respect, it should be remarked that muons are not passive spectators in the system but are perturbing the rotors in a non-trivial way. In particular, the delicate combination of the different C_3 and C_4 symmetries of the rotors and of the stators, respectively – which is ultimately allowing the preservation of motions

at cryogenic temperatures – is expected to be broken by the implantation of muons, which are adding both mass and charge in an asymmetric way. Accordingly, the detection of the onset of slower dynamical processes at temperatures higher than in the NMR measurements is reasonable. In spite of these arguments, muons still allow the estimate of an exceptionally low value for E_A , confirming the easiness of rotation of the studied molecular rotors.

We now turn to the behaviour of the other two components, $A_{f_1}^{\text{ZF}}$ and $A_{f_2}^{\text{ZF}}$. Their amplitudes sum up to $\sim 20\%$ – 25% of the maximum instrumental asymmetry, suggesting that they are associated to those muons not forming muonium. In the LT regime, both these components give rise to characteristic oscillations in ZF – see Fig. 2. Although we can use Eq. 1 at all the temperature values, we notice that better fitting results are obtained for $T \lesssim 4$ K by referring to the so-called H- μ state – see Fig. SI6. In fact, in the absence of long-range ordered magnetism in the investigated material, well-defined oscillations in the ZF- μ SR spectra are usually caused by the generation of quantum entangled states between the magnetic moments of the muons and the nuclei.^{33–35} Particularly relevant for our study is the H- μ state, where the entangled state is between the spin-1/2 nuclear spin of an hydrogen and the muon spin ($s_\mu = 1/2$). As discussed in the supporting information, our data are consistent with the generation of two distinct H- μ states well within the LT regime. Based on the conventional static model (see Eq. SI11) we quantify precisely $r_1 = 1.46 \pm 0.02$ Å and $r_2 = 0.860 \pm 0.015$ Å for the characteristic muon-hydrogen distances in the two entangled states. We stress that the strong coupling with H nuclei identifies two muon thermalization sites directly onto the rotor unambiguously. Indeed, the other nuclear species in the investigated MOF are characterized by vanishing magnetic moments and hydrogens are only located as a crown on the central plane of the bicyclo-[1.1.1]-pentane rotating moieties (see Fig. 1).

However, given the extremely low activation energy of ~ 30 cal mol⁻¹, the system still experiences rotary motion in the fast limit regime even at cryogenic temperatures and the motional averaging should reduce the apparent frequency values accordingly. Therefore,

the muon-hydrogen dipolar coupling must be averaged anisotropically because of the fast spinning of the muon on the plane perpendicular to the rotor main axis³⁶ and the distances r_1 and r_2 estimated from a static H- μ model should be renormalized as well. In the assumption of an averaging by a factor $1/2$,³⁷ the renormalized distances would be $r'_1 \simeq 1.16 \text{ \AA}$ and $r'_2 \simeq 0.68 \text{ \AA}$, which match well with muons located between the hydrogens on the rotor. Any attempt to make these arguments more quantitative should rely on a DFT modelling of the muon implantation site, which is beyond the current scopes of this study. Also, the data within the IT region show evidence of a single oscillating component but they cannot be described by the H- μ fitting function even after fixing one of the two oscillating amplitudes functions to zero. The more empirical fitting function reported in Eq. 1 gives excellent results instead, as shown in Fig. 2, with $B_1 \simeq 12 \text{ G}$. Overall, this behaviour may be due to a sudden change in the implantation sites upon increasing temperature, leading the muons to interact simultaneously with two hydrogens in a so-called H- μ -H entangled state. The functional form for the aligned H- μ -H makes the fitting quality worse, suggesting that a more subtle numerical analysis is required by assuming that the muon and the protons are not aligned.³⁴

It is remarkable that the oscillations in the ZF depolarization curves can be observed only in the LT and IT regimes. As mentioned above, the single oscillation observed in the IT region becomes progressively overdamped into an initial exponentially-decaying component upon increasing temperature into the HT region (see Fig. 2). We interpret this finding as the result of the onset of dynamical processes associated to the diffusive motion of those muons implanted on the rotors, possibly favoured by the onset of the rotary motion itself. To the best of our knowledge, this is the first report of such unusual behaviour for the evolution of H- μ coherent oscillations as a function of temperature.

A strong hint towards the dynamical origin of the initial depolarization in the HT regime arises from the behaviour of the μ SR signal under the effect of a magnetic field in LF geometry. We show selected depolarization curves for $H^{\text{LF}} = 4 \text{ kOe}$ in the inset of Fig. 4, evidencing a marked dependence on temperature. In the LT regime, we observe an almost

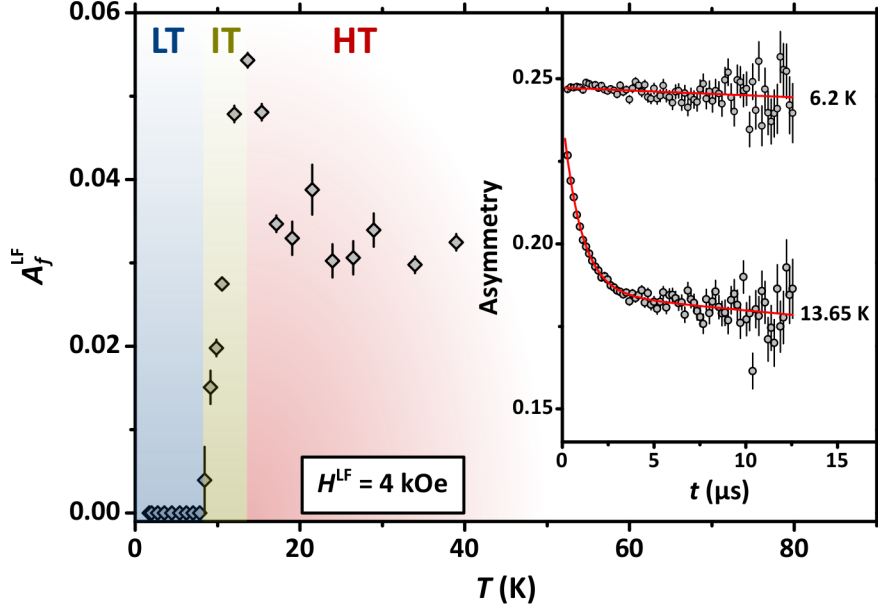


Figure 4: Main panel: temperature dependence of the amplitude of the initial exponentially-decaying component for LF- μ SR at $H^{\text{LF}} = 4 \text{ kOe}$. Inset: representative results of LF- μ SR at different temperature values and $H^{\text{LF}} = 4 \text{ kOe}$. The continuous lines are best-fitting functions according to Eq. 2.

flat depolarization with full initial asymmetry – however, we detect a clear onset of dynamical processes leading to a fast-decaying component of the muon depolarization upon increasing temperature. In the main panel of Fig. 4 we plot the temperature behaviour of A_f^{LF} showing unambiguously that dynamical processes set in upon increasing temperature within the IT region. We also notice that A_f^{LF} shows a well-defined maximum at $T \simeq 13.5 \text{ K}$ and that its amplitude is suppressed in the HT regime, leading to a markedly non-monotonic behaviour. As discussed in the supporting information (Fig. SI5), this loss of signal is not compensated by the amplitude of the long-tail component A_s^{LF} . We argue that additional processes set in within the HT region leading to a non-detectable initial depolarization because of the pulse-limited time-resolution at ISIS.

In conclusion, we used muon-spin spectroscopy at cryogenic temperatures to highlight the exceptional dynamical properties of molecular rotors bridging adjacent metal oxide nano-clusters, as engineered in a low-density metal-organic framework. Muon-spin spectroscopy provided different complementary viewpoints of the rapid rotor dynamics, even at very low

temperatures. Indeed, muons are implanted in two privileged implantation sites – one perceiving the rotor dynamics from the stator nodes, in close proximity to the rotor itself, and the other on the fast-reorienting moiety. Thanks to these favourable observatories, a rich picture of the dynamics emerged and an exceptionally low energy barrier $\sim 30 \text{ cal mol}^{-1}$ for rotations was estimated. We also detected a peculiar temperature dependence of the coherent oscillations arising from those muons implanted on the rotors and diffusing at temperatures $T \gtrsim 13 \text{ K}$, implying a non-trivial effect of rotations on the implantation itself. This study stresses the utility of muon-spin spectroscopy to identify motions in organic solids and enlarges the perspectives for its application to rotors, motors and molecular machines in the solid state.

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Supporting Information Available

We report additional details on muon spin spectroscopy and on our μSR data as supporting information. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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