Muonium diffusion dynamics in mercury oxide

J. Piroto Duarte\textsuperscript{a,*}, J.M. Gil\textsuperscript{a}, H.V. Alberto\textsuperscript{a}, R.C. Vilão\textsuperscript{a}, A. Weidinger\textsuperscript{a}, N. Ayres de Campos\textsuperscript{a}, S.F.J. Cox\textsuperscript{b,c}, J.S. Lord\textsuperscript{c}, S.P. Cottrell\textsuperscript{c}, E.A. Davis\textsuperscript{d}

\textsuperscript{a}Physics Department, University of Coimbra, P-3004-516 Coimbra, Portugal
\textsuperscript{b}Physics Department, University College London, WC1E 6BT, UK
\textsuperscript{c}ISIS Facility, Rutherford Appleton Laboratory, Chilton, Didcot, Oxon OX11 0QX, UK
\textsuperscript{d}Department of Materials Science and Metallurgy, University of Cambridge, CB2 3QZ, UK

Abstract

The diffusion dynamics of the neutral muonium state found in HgO is addressed in this work. We propose a hopping model for the diffusion, and use it to analyse time-domain µSR data. It is found that the diffusion is an incoherent quantum process, with an activation energy of 5.2(2) meV. The analysis also points to the anti-bonding site as the best-suited candidate for the muon’s localisation.

Keywords: II–VI Semiconductors; Mercury oxide; Muonium diffusion

1. Introduction

The understanding of the role of hydrogen in semiconductors has developed considerably in recent years since it was discovered that interstitial hydrogen could act as a shallow-donor impurity in compound semiconductors [1–4]. The µSR technique played a crucial role in this discovery; indeed, the very first experimental evidence was obtained with µSR experiments on cadmium sulphide and zinc oxide, where an unusually low hyperfine interaction associated with the muonium (Mu) state was identified [1,2]. Since then, systematic surveys of other compound semiconductors, mostly oxides, have enlarged the list of host systems in which the shallow-level donor muonium state is formed [5,6].

Within the context of a deep-level/shallow-level dual picture for hydrogen impurity states in semiconductors, the neutral muonium state found below 150 K in the wide-gap semiconductor mercury oxide appears somewhat as a further novelty [7,8]. Its hyperfine coupling parameters $A_{iso} = 14.93$ MHz and $D = 5.2$ MHz place it mid-way between the bond-centre muonium deep states known for long in the elemental and III–V compound semiconductors ($A_{iso}$ and $D$ ~several tens of MHz), and the anion antibonding shallow states recently discovered ($A_{iso}$ and $D$ ~hundreds of kHz). The uniqueness of this state arises from the unusual structure of HgO, shaped by broken chains running parallel to the $\alpha$-axis that coordinate the Hg and O atoms (see Fig. 1), contrasting with the much more regular tetrahedral coordination of most semiconductor systems. This makes controversial whether the neutral muonium state formed in HgO is located at the bond-centre or at the oxygen’s anti-bonding site, and no experimental evidence clearing this matter has been put forward so far.

This neutral muonium state in HgO has also been found to undergo diffusion through crystallographically equivalent positions below ionisation. The existence of diffusion dynamics has been inferred from data taken with powder samples which imply averaging of the state’s anisotropy [7]. In high-field TF geometry measurements, increasing temperature produces a gradual narrowing of the frequency powder-pattern lineshape characteristic of axially symmetric muonium; in LF repolarisation measurements, the $\Delta M = 1$ level-crossing resonance dip narrows from an asymmetric powder-shape at low temperatures to a more symmetrical resonance at higher ones.

This work reports a quantitative µSR study performed on the diffusion dynamics of muonium in HgO. Existing
and new experimental data taken on powder samples with the GPS (TF geometry) and EMU (LF geometry) spectrometers at PSI and ISIS respectively are analysed with a simplified hopping model, and its results are discussed.

2. Diffusion model

Presuming the observed muonium state has the same symmetry axis as the Hg–O bond, hopping paths through bond-centre or anti-bonding sites in adjacent chains provide the anisotropy averaging observed in Ref. [7] (Fig. 1). Careful examination shows that in each path jumps take place between positions either with the same symmetry axis, or with symmetry axis concurrent at $107.3^\circ$, the Hg–O–Hg bond angle. For the sake of simplicity, one may start by assuming that all polarisations of the muonium system (muon and electron) do not suffer any change when it jumps between sites having the same symmetry axis. This reduces the hopping problem to the reversible transition between two muonium states with the same hyperfine parameters, but different symmetry axis, at a hopping rate $A$. We note that these considerations apply to both the bond-centre and the oxygen anti-bonding paths, and that therefore the model makes intrinsically no distinction between the two possibilities. We also make clear that the hopping rate $A$ refers to jumps between positions with different symmetry axis, i.e., every second position in the paths; the real hop rate will be the double of $A$.

The time evolution of the muon’s polarisation in LF and TF geometries was computed using the equations of motion for all polarisations of two muonium systems with different hyperfine interaction tensors undergoing a reversible transition at a rate $A$ [9]. The procedure followed is a generalisation of the Wangsness–Bloch equations for a single muonium system to the two system case, adopting the remarks described in Ref. [10]. The two hyperfine tensors were constrained to have the same hyperfine parameters $A_{iso} = 14.93$ MHz and $D = 5.2$ MHz, and symmetry axis concurrent at $107.3^\circ$. Powder averaging was performed numerically with the (also numerical) solutions of the two-muonium system equations. A phenomenological electronic depolarisation rate $v$ was included as well, in order to model the nuclear hyperfine interaction with the surrounding spin-carrying Hg nuclei (17% $S = \frac{1}{2}$, 13% $S = \frac{3}{2}$).

Simulations performed with this model were found to reproduce the main features observed in the diffusion data above 10 K, namely the expected onset of the powder-distribution narrowing in high TF and of the narrowing of the LF resonance dip to a symmetric shape at a jump rate near the value of the anisotropy parameter. This study also showed that the high-TF and LF repolarisation data should exhibit a very low sensitivity to the hop rate $A$ outside that region. It was found as well that the highest sensitivity to $A$ is achieved with data collected in the so-called Mu* LF magic field [9,10], at which the amount of information regarding $A$ in the time spectra is maximal thanks to the non-existence of damping of the $f_{12}$ precession by the powder distribution.

3. Results and discussion

Most of the quantitative analysis with the diffusion model was performed on magic field time-spectrum data collected at the EMU spectrometer. Fig. 2 shows an Arrhenius plot of the temperature dependence extracted in that way for the Mu hop rate (double of $A$, see Section 2). Between 6 and 30 K, it is in good agreement with the simple small-polaron model for incoherent quantum diffusion of light interstitials [11,12], bearing an activation energy of $E_a = 5.2(2)$ meV and transition matrix element $J = 0.065(3)$ meV. Such small activation energy confirms the quantum nature of the diffusion process, discarding classical “over-the-barrier” hopping. The transition matrix element $J$ is also within the expected range of values for insulators and semiconductors at low temperatures [12]. At 30 K, the hop rate peaks; muonium is highly mobile at that temperature, tunneling about 800 times during its 2.2 $\mu$s lifetime. Above 30 K, the hop rate decreases, in what appears to be a change of diffusion regime. Trapping at a defect is not the mechanism responsible for this decrease, since the average length traveled by muonium at 30 K considering a one-dimensional random walk is only 60 Å, a
distance too small to allow muonium finding any impurity if one accounts for the high purity of the samples. Trapping at a grain boundary is ruled out in a similar way.

Insight about the muon site may also be derived from the magic-field analysis with the diffusion model. The electronic depolarisation rate $v$ was seen to increase with temperature from zero to $0.25(10)$ MHz at 10 K, decreasing thereof again to zero. Since $v$ is expected to model the nuclear hyperfine interaction with the Hg nuclei, its peak value may be taken as a guess for that interaction’s coupling. A rough estimate of the spin-carrying Hg nuclei to muon distance assuming a 1s type electronic wave function for the muonium computed with that coupling amounts to about 6 Å, a value far too large for the bond-centre site hypothesis, even taking into account that only one third of the Hg nuclei have spin. This is in agreement with recent theoretical results which find electrostatic stable positions for the muon at sites anti-bonding to the oxygen atom [13].

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