Anisotropic superconducting properties of aligned MgB$_2$ crystallites

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Samples of aligned MgB$_2$ crystallites have been prepared, allowing for the first time the direct identification of an upper critical field anisotropy $H_{c2}^{ab}/H_{c2}^{c} = \xi_{ab}/\xi_{c} \approx 1.7$, with $\xi_{ab} \approx 70$ Å, $\xi_{o,c} \approx 40$ Å, and a mass anisotropy ratio $m_{ab}/m_{c} \approx 0.3$. A ferromagnetic background signal was identified, possibly related to the raw materials purity.

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The recent discovery of superconductivity at 39 K in Magnesium Diboride (MgB$_2$) has brought new excitement to the area of basic and applied research on superconducting materials. The observation of an isotope effect, a BCS-type energy gap measured by Scanning Tunneling Spectroscopy, as well as band structure studies, point to a phonon-mediated superconductivity in MgB$_2$. Some reports have suggested that MgB$_2$ has an isotropic (or 3D) behavior, based on measurements done in polycrystalline samples. However, other studies have also discussed its possible anisotropic nature. The relatively high values reported for the critical current density $J_c$ are possibly indicating the absence of weak link problems, which are well known in the high-$T_c$ materials. While polycrystalline MgB$_2$ is very easy to grow and is a readily available reagent, good-sized single crystals of this material have not yet been reported, and their development promises to be a greater challenge.

Here we present results from samples of aligned MgB$_2$ crystallites that establish the anisotropy of the upper critical field ($H_{c2}$), thus implying an anisotropic character for other superconducting properties, e.g., the energy gap, coherence length ($\xi$), field penetration depth ($\lambda$), and $J_c$.

In this work, a weakly sintered sample of MgB$_2$ was prepared, starting with a stoichiometric mixture of 99.5 at% pure Boron and 99.8 at% pure Magnesium, both in chips form (Johnson Matthey Electronics). The loose mixture was sealed in a Ta tube under Ar atmosphere, which was then encapsulated in a quartz ampoule and put into the furnace. The compound formation was processed by initially holding the furnace temperature at 1200°C for 1 hour, followed by a decrease to 700°C (10°C/h), then to 600°C (2°C/h), and finally to room temperature at a rate of 100°C/h. The weakly sintered product was easily crushed and milled employing mortar and pestle. Using a stereomicroscope we could observe a very uniform powder consisting mainly of shiny crystallites, with aspect ratios ranging from 2 to 5. This is mainly due to the main surface size distribution ranging from 5 to 40 µm for the larger linear dimension, since the crystallites’ thickness is very regular, around 2 µm. The powder was then sieved into a range of particle sizes between 5 - 20 µm, which allows the crystallites fraction to be maximized to almost 100%. Small amounts of the powder were then patiently spread on both sides of a small piece of paper, producing an almost perfect alignment of the crystallites, as shown in the SEM picture in the upper part of Fig. 1. The lower part of this figure shows an X-ray diffraction pattern ($\theta - 2\theta$ scan) from a sample of the crystallite-painted paper, displaying only the (001) and (002) reflections coming from the MgB$_2$ phase. A lattice parameter $c = 3.518 \pm 0.008$ Å was evaluated from these two peak positions. The two small impurity peaks marked with asterisks were indexed as SiO$_2$. The inset of Fig. 1 shows a rocking curve ($\omega$ scan) for the (002) peak that reveals an angular spread around 4.6 degrees, associated with a small misalignment of the crystallites $c$ axis.

Electron microprobe analysis done on four different areas between the MgB$_2$ crystallites, revealed the following average concentration (in at%) of elements: O (62.9), C (22.2), Ca (9.48), Si (1.48), Mg (1.44), Al (1.37), K (0.09), Fe (0.50), Cr (0.21), Ni (0.09). The first eight elements in this list were found also in the composition analysis made on the same type of paper used (Canson, ref. 4567-114). Microprobe analysis done also on the initial Mg and B revealed a few small precipitates, smaller than 10 µm and containing up to 8 at% Fe, only in the Mg chips. This confirms the expectation of Fe being a common impurity in commercial Mg and sets a general concern on its possible effects. The average composition found on top of several crystallites, normalized to the whole MgB$_2$ formula unit, was: Mg (30.80), O (2.20), Ca (0.17), Si (0.07), Fe (0.06). Although Boron contributes with a fraction of 66.6 at% it does not show-up in the microprobe analysis because it is too light. The contaminants found on top of the crystallites most possibly came from a surface contamination caused by the alignment technique, which required vigorous rubbing on top of the powder, using a steel tweezers tip to spread the crystallites uniformly. This is corroborated by a further analysis done on top of several as-grown crystallites, which detected only Mg and a small amount of O (possibly from MgO). This result is consistent with the very small solid solubility limit of about 0.004 at% Fe in Mg, which is known to occur at the solidification temperature of 650 °C. The inter-crystallite type of rubbish shown in Fig. 1 is attributed mainly to the paper abrasion, which produces a varied distribution of irregular grains of paper.
fragments. In order to characterize the superconducting and magnetic properties of the aligned crystallites, we mounted several samples consisting of a pile of 5 small squares (3 × 3 mm²) cut from the crystal-like-painted paper and glued with Araldite resin. Each one of these samples contains a number of crystallites estimated to be around 6.5 × 10⁵, totaling an effective volume of 0.005 mm³, which is reasonably close to 0.060 mm³ that was evaluated from the expected slope of −1/4π for the diamagnetic shielding at \( H ≈ 0 \).

Figure 2 shows the anisotropic signature of the \( H_{c2}(T) \) line in the field interval \( 0 \leq H \leq 40 \text{ kOe} \). The values were taken from the transition onset of the real component (\( χ' \)) of ac susceptibility, measured using a PPMS-9T machine (Quantum Design), with an excitation field of amplitude 1 Oe and frequency 5 kHz. The inset shows an enlarged view of the \( χ'(T) \) curves for H parallel (solid symbols) and perpendicular (open symbols) to the sample c axis. The \( χ'(T) \) as well as the \( M(T) \) (inset of Fig. 3) measurements, for \( H = 10 \text{ Oe} \), show sharp transitions at the same critical temperature \( T_c = 39.2 \text{ K} \). The dashed lines connecting points in Figs. 2 - 4 are only guides to the eyes. Typically, some of the published data on the temperature dependence of \( H_{c2}(T) \) agree with our result for \( H_{c2}(T) // ab \). As an example, the data from Ref. 14 is plotted in Fig. 2 as stars. This could simply mean that in polycrystalline samples the transitions are broadened, showing the onset at the highest temperature that corresponds to the highest critical field available, which is \( H_{c2}(T) // ab \).

The ratio \( η = H_{ab}^{c2}/H_{c2}^{c} \), between the upper critical field when \( H \) is applied parallel to the \( ab \) plane, and when it is along the c direction, was evaluated at different temperatures, producing \( η = 1.73 \pm 0.03 \). Using the Ginzburg-Landau mean field expression \( ξ(T) = \xi_0(1 - T/T_c)^{-1/2} \) and the results for anisotropic situations, \( \xi_{ab}^{c2}(T) = \phi_0 / (2 \pi \xi_{ab}^{c2}) \) and \( H_{ab}^{c2}/H_{c2}^{c} = 1/ξ_c \), where \( \phi_0 = 0.207 \times 10^{-7} \text{ G cm}^{-2} \) is the quantum of flux and \( \xi^2 = m_{ab}/m_c \) is the mass anisotropy ratio, we find that \( \phi_{ab}/\xi_c = \xi_{ab}^{c2}/ξ_c(T) = \eta ⇐ 1.73 \) and \( ξ^2 ≈ 0.3 \). Since at \( T = 27 \text{ K} \) we have \( H_{c2}^{c} = 20 \text{ kOe} \), this implies that \( \xi_{o,ab} ≈ 70 \text{ Å} \) and \( \xi_{o,c} ≈ 40 \text{ Å} \). The mass anisotropy ratio of MgB₂ thus corresponds to a relatively small anisotropy when compared to the highly anisotropic high-Tc cuprates like YBCO (\( \xi^2 ≈ 0.04 \)) and BSCCO (\( \xi^2 ≈ 10^{-4} \)). We do not expect that a likely very small bulk contamination of the crystallites could eventually change their anisotropy values. In fact our careful composition analysis have indicated that almost all contaminants are located in the region between the crystallites, thus having a negligible chance to affect the underlying mechanism of the superconducting condensation.

The magnetization curves \( M(T) \) and \( M(H) \), displayed in Figs. 3 and 4, were measured using a SQUID magnetometer (Quantum Design, model MPMS-5). The \( M(H) \) curves (\( T = 5 \text{ K} \)) shown in Fig. 3 are intriguing in the region \(-1 \leq H \leq 1 \text{ kOe} \), where the maximum shielding and first field penetration (in the initial virgin state) occur. For \( |H| > 1 \text{ kOe} \) the hysteretic curves in both field directions look very similar. However, for \( |H| > 40 \text{ kOe} \) (not shown here) the magnetization difference between the up and down curves (\( ΔM \)) becomes smaller than the noise. Large fluctuations of the magnetic moment were consistently observed in this field region, for 3 different samples and temperatures (\( T = 5, 10, 20 \text{ K} \)), possibly associated with the high open ratio and the fast drop of \( J_c \) occurring at high fields. The signature of the ferromagnetic hysteresis loop measured at \( T = 45 \text{ K} \), mainly attributed to the presence of Fe, Cr and Ni in the inter-crystallite region. The inset displays an enlarged view close to \( H = 0 \) indicating that demagnetization effects are also observed for the \( H // ab \) and \( H // c \) orientations. In a recent detailed study, the occurrence of Fe contamination has already been identified, through measurements of MgB₂ samples made from commercial powder supplied by a different company.

In view of the superimposed ferromagnetic signal in the magnetization curves, we found to be not reliable to discuss the expected anisotropy in \( J_c \propto \xi_{ab}/ξ_c \), which could be determined using the Bean model. A rough estimate for both field orientations gives \( J_c \propto 10^6 \text{ A/cm}^2 \) at \( H = 1.5 \text{ kOe} \) and \( T = 5 \text{ K} \) (Fig. 3). This calculation neglects the small influence of the ferromagnetic hysteresis and considers the average crystallite geometry as described before. However, an anisotropy between \( J_c(H//c) \) and \( J_c(H//ab) \) should be expected. Indeed, independently of the different regime of vortex pinning, \( J_c \) is predicted to be proportional to \( ξ^2 \), leading to \( J_c(H//c) / J_c(H//ab) ≈ (ξ_{ab}/ξ_c) ≈ H_{ab}^{c2}/H_{c2}^{c} \).

A final cautionary observation has to be addressed to the possibility that surface superconductivity could also be occurring for \( H // ab \), since coincidently the surface nucleation field is \( H_{c3} ≈ 1.7 H_{c2} \). However, we have made several careful measurements of \( M(H) \) and \( χ'(H) \), as well as \( M(T) \) and \( χ'(T) \), around the onset of transition, and no signature of a surface nucleation field was found. This is consistent with the fact that our \( H_{c2}^{c}(T) \) line agrees with several reported \( H_{c2}^{c}(T) \) lines measured in polycrystalline MgB₂, which certainly did not comply the boundary conditions required for surface nucleation in the \( ab \) planes, i.e. \( H // ab \).

In conclusion, we have prepared samples of aligned MgB₂ crystallites that allowed, for the first time, the identification of an anisotropy for the upper critical field given by \( H_{ab}^{c2}/H_{c2}^{c} = 1.73 \), implying an anisotropy of the coherence length \( ξ_{ab}/ξ_c ≈ 1.73 \) and a mass anisotropy ratio \( m_{ab}/m_c ≈ 0.3 \). This could be considered a mild anisotropy when compared to the values found for the high-Tc materials \( m_{ab}/m_c ≈ 0.04 \). The influence of contaminants is requiring further work, aimed at a more complete and reliable characterization of the MgB₂ intrinsic properties. Naturally the production of a good-sized single crystal of MgB₂ is also highly desirable.

Note added: Since this manuscript was submitted two papers have appeared showing results consistent
with our anisotropy data.

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FIGURE CAPTIONS

FIG. 1. Top: SEM picture showing the well aligned crystallites and inter-crystallite material. Bottom: X-ray diffraction pattern showing only the (001) and (002) peaks of MgB$_2$, plus two spurious peaks indexed as SiO$_2$. Inset: rocking curve (ω scan) for the (002) peak, showing an angular spread of about 4.6 degrees along the crystallites c axis.

FIG. 2. Upper critical field $H_{c2}$ vs. Temperature phase diagram, for both sample orientations. The stars represent the $H_{c2}$ vs. $T$ line from Ref. 14. The inset shows the real component $\chi'$ of the ac susceptibility vs. temperature, measured at several dc fields for both orientations. Open symbols are for the $H//ab$ curves and solid symbols for $H//c$.

FIG. 3. Magnetization loops at 5 K for both sample orientations, showing a superconducting hysteresis on a ferromagnetic background. The inset shows a dc magnetization vs. temperature curve at 10 Oe, showing a sharp transition at 39.2 K and $\sim 70$ % recovery of diamagnetism for the FCC curve.

FIG. 4. Magnetization loops at 45 K (above $T_c$) for both sample orientations, showing the ferromagnetic behavior of our sample. The inset shows the hysteretic behavior at low fields.
FIG. 1

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H$_{c2}$ (kOe) vs. T (K)

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Fig. 2
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**FIG. 3**
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FIG. 4