Effect of ball milling on the local magnetic flux distribution and microstructure of in situ Fe/MgB$_2$ conductors

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Abstract:

The effects of the precursor milling energy on the superconducting and microstructural homogeneity of Fe-sheathed MgB$_2$ monocore wires and tapes have been investigated. The conductors were produced by the powder-in-tube method using drawing and rolling deformation methods and in situ reaction. Combined magneto-optical, structural and magnetization measurements were performed on MgB$_2$ superconducting tapes prepared from differently milled precursors revealing hidden mechanisms controlling their critical current density. Local degradation of the superconducting properties observed in some analysed conductors has been identified and correlated with microstructural observations.
1. Introduction

MgB₂ conductors have been the subject of intense research in the last years to improve the superconducting properties and optimize the manufacturing process towards the development of homogeneous and long conductors with the performance required for power applications. With the critical temperature $T_c \sim 39$ K, this superconductor allows operating temperatures about 20 K, which implies enhanced thermal stability of superconducting devices and lower operating cost than for systems based on low temperature superconductors, such as NbTi and Nb₃Sn, typically working at temperatures below 5 K. On the other hand, compared with available high $T_c$ superconductors based on YBa₂Cu₃O₇−x (YBCO) and Bi₂Sr₂Ca₂Cu₃O₁₂+x (Bi-2223), the low manufacturing cost and cheap constituent materials of MgB₂ conductors can open market niches in applications such as Magnetic Resonance Imaging, MRI [1-4], Superconducting Magnetic Energy Storage, SMES [5] or superconducting wind turbines [6].

Highly uniform and technologically reproducible critical current density, $J_c$, in MgB₂ conductors is compulsory for advanced applications. To achieve this goal, a homogeneous microstructure and phase composition distribution along the superconducting wire is required. Grain refinement of MgB₂ would improve homogeneity and increase the critical current of the material due to larger density of grain boundaries, which would provide additional pinning [7,8]. Very fine grained microstructure has been widely searched by mechanical ball-milling of different precursor powders: MgB₂ [9], Mg + B [10-14], B [15-18], or Mg [12], to prepare bulks and wires with the ex situ and in situ reaction methods.

By increasing the milling energy transferred to the precursor, the irreversibility and upper critical fields are also increased, which causes a significant enhancement of $J_c$ in the high field region [12,13,14,16,18]. This is a result of a poorer crystallinity of superconducting grains that induces lattice strain but also causes a small reduction of $T_c$ [15,18,19]. Häßler et al [13] have correlated the observed $J_c(H)$ enhancement at low temperatures and high fields with the milling energy, which depends on the rotation speed and milling time, and found a transition from grain size reduction to mechanical alloying at milling energy of about $10^8$ J kg⁻¹.

On the other hand, milling procedures may also produce some undesirable effects such as a decrease of $J_c$ values at low magnetic fields [14,15,16,18], higher oxygen content [16,18] and, in some cases, a degradation of the deformation properties during wire manufacture [13,14]. In order to further explore the reasons of such a decrease in $J_c$ in the low field range, in this contribution we have analysed the local magnetic flux distribution inside MgB₂ conductors made from different energy milled precursors using magneto-optical (MO) imaging. Local superconducting properties have been correlated with magnetization measurements and microstructural observations by scanning electron microscopy (SEM).
2. Experimental details

Monocore Fe-sheathed MgB$_2$ wires were prepared by the powder-in-tube (PIT) method using the in situ reaction technique. The starting materials were amorphous B (99%, New Metal & Chemicals Ltd, mean particle size lower than 1 μm) and Mg (Goodfellow 99.8% purity with grain size lower than 250 μm). Mg and B powders in stoichiometric proportions 1:2 were mixed in a vibratory mill Retsch MM 200 for 30 minutes. Afterwards, some of these precursor powders were milled in Ar atmosphere using a planetary ball mill (Retsch PM 100), with sun disk of radius $r_p = 7$ cm and tungsten carbide (WC) jar with inner radius $r_v = 5.25$ cm. The ball-to-powder mass ratio, $\beta$, was kept at $\sim 36$. During the milling process, one-minute pauses and change in rotation direction every three minutes were chosen to avoid local heating and formation of MgB$_2$ by mechanical alloying. The effective ball milling time ($t$) and rotation speed ($\omega_p$) are given in Table 1 for each analysed wire. Three milling conditions were used: M1 (200 rpm and $t = 1.5$ h), M2 (400 rpm and $t = 3$ h) and M3 (400 rpm and effective milling time of 16.5 h).

Fe tubes (99.5%) of 5 mm outer diameter and 0.25 mm wall thickness were hand-filled with the different precursor powders and sealed at both ends. The diameter was reduced by drawing in consecutive steps down to the final wire diameter of 1.1 mm. An intermediate annealing during drawing was performed to reduce the Fe sheath’s work hardening (550 °C for 1 h in Ar). Since MO imaging requires flat surface, some pieces of these wires were rolled to tapes (of $\sim 2$ mm width and $\sim 0.4$ mm thickness) before the final annealing. Finally, pieces of length $\sim 10-13$ cm were sealed to prevent oxidation, and heat treated at 670 °C for 5 h in vacuum (0.1 mbar) to form the superconducting MgB$_2$ phase. The main characteristics of the analysed wires, including a reference sample with unmilled precursor (M0), are collected in Table 1.

**Table 1.** Manufacturing characteristics and properties of the analysed Fe-MgB$_2$ wires and tapes: $t$ is the effective ball milling time, $\omega_p$ is the mill rotation speed and $E_t/m$ is the estimated energy per unit mass transferred to the precursor (eq. 1). $T_c$ and $\Delta T_c$ are the critical temperature and the transition width, respectively, as derived from $\chi_{ac}(T)$ measurements of the wires.

<table>
<thead>
<tr>
<th>Wire</th>
<th>Tape</th>
<th>Ball milling</th>
<th>$T_c$ (K)</th>
<th>$\Delta T_c$ (K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>W-M0</td>
<td>T-M0</td>
<td>0</td>
<td>0</td>
<td>37.5</td>
</tr>
<tr>
<td>W-M1</td>
<td>T-M1</td>
<td>1.5</td>
<td>200</td>
<td>$1.2 \times 10^6$</td>
</tr>
<tr>
<td>W-M2</td>
<td>T-M2</td>
<td>3.0</td>
<td>400</td>
<td>$2 \times 10^7$</td>
</tr>
<tr>
<td>W-M3</td>
<td>T-M3</td>
<td>16.5</td>
<td>400</td>
<td>$1 \times 10^8$</td>
</tr>
</tbody>
</table>

The microstructure and phase composition of the final wires and tapes were analysed by field-emission scanning electron microscopy (FE-SEM, Carl Zeiss
MERLIN), with 15 kV accelerating voltage and probe current of 600 pA. The phase composition was analysed using the X-ray diffraction (XRD) patterns collected at room temperature on a RIGAKU D-max /2500 X-ray diffractometer using Cu Kα radiation.

Vibrating sample magnetometers (VSM, Quantum Design PPMS-9T and PPMS-14T) were used to measure isothermal magnetic hysteresis loops \( M(H) \). AC magnetic susceptibility as a function of temperature, \( \chi_{ac}(T) \), measured with a SQUID-based Quantum Design MPMS-5T, was used to estimate the critical temperature \( T_c \), defined as the diamagnetism onset, and the transition width \( \Delta T_c \), defined as the change of \( \chi_{ac} \) in-phase component (\( \chi' \)) from 10% to 90% of its value at 5 K. For this magnetic characterisation, the iron sheath was removed by mechanical polishing because its strong ferromagnetic signal hinders the superconductor’s contribution analysis. AC and DC fields were applied perpendicular to the axis of ~ 4-5 mm long wires. Electrical DC transport measurements in perpendicular DC magnetic fields were performed in some wires with the 6-cm long wire immersed in liquid helium. The critical current densities, \( J_c \) were determined by the standard four-probe method using the 1 μV/cm criterion.

Magnetic flux distribution was visualised in a linear polarised light using magneto-optical (MO) imaging technique based on the rotation of the light polarization plane in presence of a magnetic field (the Faraday effect) [20,21]. As field sensors we used in-plane magnetized bismuth-substituted iron garnet sensor films [21], which provide strong rotation in a weak magnetic field and give a desirable sensitivity and spatial resolution for monitoring local magnetic field on the surface of the films. The MO indicator film is placed directly on top of the polished surface of the sample and the light is reflected from the mirror on the bottom of the film before going into the analyser of the optical system set above the sample holder. The brightness of the images obtained with crossed polarizers represents the absolute value of the magnitude of the z-component of local magnetic field in the sample.

We used a polarized-light microscope and a charge coupled device (CCD) to capture and process the digital images obtained in the experiments. The microscope consists of a stabilized light source, a polariser, an analyser and optical components to project the image plane into the CCD camera with various magnifications. All optical components, especially the objective lenses, have small Verdet constants [20] to not depolarize the light beam. The magneto-optical measurements were carried out in transmission mode with sensitivity defined by the thickness of the magneto-optical layer the light is going through.

3. Results and discussion

3.1 Effect of ball milling on the global properties of MgB₂ conductors

The energy per unit mass transferred to the powder by ball milling, \( E/l/m \), is a useful parameter to analyse the milling effects. There are different approaches to
determine $E_t/m$. In this work we have used the formula developed in [13], which depends explicitly on the milling parameters defined in section 2:

$$\frac{E_t}{m} = c\beta \left(\omega_p r_p\right)^3 \frac{1}{r_v} t,$$

where $c$ is a dimensionless constant on the order of 0.1, $\beta$ the ball-to-powder mass ratio, $\omega_p$ the rotation speed, $r_p$ the sun disk radius, $r_v$ the jar inner radius and $t$ the effective ball milling time. It must be noted that an unknown fraction of this energy is lost to heat, to plastic deformation of the powders or even to form MgB$_2$ by mechanical alloying. According to Eq. (1), the used milling conditions M1, M2 and M3 described above correspond to energy values $E_t/m \approx 1.2 \times 10^6$, $2 \times 10^7$ and $1 \times 10^8$ J kg$^{-1}$, respectively.

XRD pattern of the precursor powder with milling conditions M3 indicates the presence of already reacted MgB$_2$ (mechanical alloying), as shown in Figure 1, in agreement with [13], whereas for the other milling conditions only the peaks corresponding to Mg are visible. Boron is in amorphous phase and therefore it is not visible in the XRD patterns. After the heat treatment of the final wire, the main phase in all wires corresponds to the superconducting MgB$_2$. The main secondary phase is MgO, and also WC in the wires made from milled powders.

![XRD pattern of the precursor powder with milling conditions M3.](image)

**Figure 1** XRD pattern of the precursor powder with milling conditions M3.

The precursor milling conditions affect $T_c$ and $J_c$ values. As shown in Table 1, $T_c$ decreases when the milling energy increases, but the transition remains always sharp, with $\Delta T_c$ values in a range between 0.5 and 1 K for all wires. Figure 2 displays $J_c(H)$ at 5 and 20 K of wires made with unmilled and milled powders. The values were obtained from the width of the hysteresis loop, $\Delta M(H)$, measured at constant temperature, using the equation: [22]

$$J_c(H) = \frac{3\pi}{8} \frac{\Delta M(H)}{R},$$

(2)
which is valid for wires of radius $R$ and length $L$, $L >> R$, under magnetic fields perpendicular to their axis. Magnetic flux jumps limit the magnetization values at low magnetic fields (below 2 - 3T at 5 K for all wires and below 1.4 T at 20 K for W-M0) leading to underestimations of $J_c$ values in this field range, which are not displayed. The two opposite effects of ball milling on $J_c$ are clearly seen in Figure 2. As mentioned before, on the one hand, higher energy milling enhances vortex pinning at high fields, so that the highest $J_c$’s at 5 K and $\mu_0H > 5$ T correspond to W-M3. On the other hand, at low fields and higher temperatures (20 K and $\mu_0H < 3$ T), $J_c$ decreases with increasing milling energy, which indicates poorer grain connectivity and/or larger amounts of microstructural defects than in the unmilled sample. Measurements of transport $J_c$ of wires W-M0 and W-M1, gave values of $6 \times 10^7$ Am$^{-2}$ at 4.2 K and 8 T for W-M1 and $4-5 \times 10^7$ Am$^{-2}$ for W-M0 (for different parts of the same wire).

![Figure 2](image.png)

**Figure 2** Magnetic field dependence of the inductive critical current density at 5 and 20 K for wires made from different ball milled precursors as specified in Table 1.

### 3.2 Microstructural and Magneto-Optical analysis

Magneto-optical imaging was performed to evaluate the homogeneity of the superconducting properties across and along the conductors. As mentioned previously, this analysis requires flat surfaces, so it has been used for characterising the tapes. One of the tapes’ faces was polished to remove the Fe sheath, so that there is not magnetic shielding by the ferromagnetic Fe sheath under perpendicular fields and gives direct access to the superconducting core both in MO experiments and microstructural SEM analysis.
Figure 3 shows SEM and MO images of tapes made from powders with increasing milling energies from top to bottom. SEM and MO images correspond to the same area of the sample, except in Fig. 3(d) where different areas of the same tape were analysed due to technical reasons. SEM images were taken with the secondary electrons (SE) detector to analyse the topographical differences among the samples. MO images were taken after zero-field cooling (ZFC) the sample from a temperature above $T_c$ down to 20 K, then applying a field of 85 mT perpendicular to the sample surface and finally reducing the field to zero, to show the remnant flux density. It should be noted that the length of the tapes analysed by MO imaging is about five times larger than the core width ($\sim$10 and $\sim$2 mm, respectively) in all samples.

**Figure 3** SEM (left) and corresponding MO images (right) of different tapes made from powder with increasing milling energies from top to bottom. SEM images were taken with the secondary electrons detector. MO images were taken after ZFC to 20 K, then applying a field of 85 mT perpendicular to the tape and finally removing the field. SEM and MO images correspond to the same area, except in (d) where images of different areas of the same tape were analysed. The inset in (d) shows a magnified image of the core near the sheath. The same magnification was used for both SEM and MO images.

As reported previously [14], upon increasing the milling energy, the microstructure of the superconducting cores changes gradually, forming smaller grains and voids, but also producing variations on the voids orientation and showing new types
of defects. The SEM images of the tapes shown in Figure 3 follow the same trend. In the specimen made from unmilled powders (Figure 3(a)-left), elongated voids parallel to the tape axis, with size of several hundred microns, are observed. The reason for such a microstructure is that Mg grains are deformed into elongated shapes during drawing and rolling due to their higher plasticity than boron. During the subsequent heat treatment, the Mg diffusion into nearby boron grains and the higher density of MgB$_2$ compared to the starting powders produce elongated voids. The shape of the magnetic flux front obtained by MO imaging, Figure 3(a)-right, follows this porous structure. It must be remarked that the orientation and shape of the voids in this sample facilitates the magnetic flux penetration from both left and right ends towards the centre, compared to the other analysed samples.

The SEM images shown in Figure 3 indicate that the use of precursors with increasing milling energy decreases gradually the porosity of the final tape. The MO image of the tape made with low-energy milled precursors, T-M1, Figure 3(b)-right, shows good and homogeneous superconducting properties. Note that during polishing of this tape, some Fe particles from the tape’s sheath were dragged towards the superconducting core surface (light grey particles in SEM, Figure 3(b)-left).

Tape T-M2, Figure 3(c)-left, has a dense core but also some microstructural defects, such as pores in the Fe sheath (seen in the upper-side of the image), which affect significantly the superconducting properties of the core around, as it is seen in Fig 3(c)-right. This sample also has a region with lower local $J_c$ (deeper penetration of the magnetic flux) in the upper-right part of the image, which, again, coincides with a microstructural defect.

Finally, the tape made from the highest energy milled precursor (T-M3) has cracks perpendicular to the long direction of the tape, as clearly seen in Figure 3(d)-left. MO image (Figure 3(d)-right) shows also deep magnetic flux penetration along these cracks that cross the superconducting core and strongly decrease $J_c$ in this tape. During mechanical deformation, the fine-milled precursor forms highly dense blocks separated by less-denser packed powders or even cracks that cross the tape from side to side perpendicular to it. This forms the brick-like structures that cannot be restored by the final heat treatment. Moreover, SEM images show that the core is very dense in the centre of this tape, while at the edges (upper and lower areas in Figure 3(d)-left), microstructural defects such as cracks and voids are present, which produce magnetic flux patterns with ripple-like shape, as shown in Figure 3(d)-right. It must be mentioned that a structure of flux penetration along cracks running perpendicular to the tape axis were also observed by MO imaging on Bi-2223 tapes after bending and straightening the samples below the critical bending diameter, which causes $J_c$ degradation due to crack formation [23]. However, the transversal cracks observed in tape T-M3 were produced during rolling not from bending the sample.
Since the reasons of the observed deterioration of superconducting properties in the sample T-M2 were unknown, further MO experiments and SEM analysis were performed, comparing it with tape T-M1, which shows the best behaviour among the analysed tapes. With this purpose, MO images were recorded with the following thermal and magnetic history: the tape was field-cooled (FC) from $T>T_c$ in a perpendicular field of 17 mT down to 20 K. Next, after stabilizing the temperature, the field was decreased to zero. Subsequently, the field amplitude was increased again but in the opposite direction to -85 mT, and finally returned back to zero. The results for tapes T-M2 and T-M1 are shown in Figures 4 and 5, respectively. The $M(H)$ curves for similar magnetic history, but measured in different pieces of the same tapes, are also shown. MO images taken in a field-cooled sample following this specific magnetic field protocol highlights the distortions in the magnetic pattern so that it is helpful to reveal inhomogeneities in the superconductor.

Figure 4 shows MO images of tape T-M2 taken at different magnetic fields during this measurement. The image (not shown) taken just after field cooling the sample to 20 K in 17 mT, displays a uniform grey background. Upon decreasing the field to zero (b) a magnetic flux inhomogeneity (area marked by an arrow) becomes visible in the MO image, and gets clearer when further increasing the field amplitude in the opposite direction, as shown in images (c) and (d). Note that at the magnetic field of -85 mT, shown in (d), the shape of the penetration front is similar to that in Figure 3(c). When slightly decreasing the amplitude of the field again to -76.5 mT, image (e), some new features in the penetration field front appear suddenly in the upper part of the image (marked by arrows), and also some “bubble like shape” magnetic inhomogeneities are formed in the central part of the sample. It must be noted that the initial round inhomogeneity is always observed in the same position in all measurement runs performed both at 5 and 20 K. While the bubble-like feature, always suddenly formed in the same area, is not always of the same structure. This behaviour suggests the existence of poor superconducting links between regions in this part of the sample. Finally, MO image (f) shows the trapped field, which has similar pattern as in (e) but with the expected dark-bright-dark profile revealing the existence of flux-antiflux areas due to hysteresis and the performed thermal-magnetic history. It must be remarked that the left part of this sample shows good homogeneity and better superconducting properties than the centre and right part of the sample.
Figure 4 (a) Magnetization values, $M(H)$, and (b)-(f) MO images of the tape T-M2 at different fields after FC in 17 mT to 20 K, then decreasing the field to -85 mT and finally removing the field. The images correspond to: (b) 0 mT, (c) -17 mT, (d) -85 mT, (e) -76.5 mT and (f) 0 mT, as marked in (a). The magnetic field is applied perpendicular to the tape. The sample of MO images is the same as in Figure 3(c), while $M(H)$ was measured in a different piece.
Figure 5 (a) Magnetization values, $M(H)$, and (b)-(c) MO images of the tape T-M1 in zero-field of the upper and lower branch, respectively, as marked in (a). Similar thermal and magnetic history as in Figure 4 was performed: the sample was FC in 17 mT to 20 K, the field was then decreased to -85 mT and finally returned back to zero. The sample of MO images is the same as in Figure 3(b), while $M(H)$ was measured in a different piece.

For comparison, Figure 5 shows MO images of the tape T-M1 during a similar measurement run as in Figure 4. It must be emphasised that the upper and lower branches of $M(H)$ here are very similar, indicating small penetration of the magnetic field and therefore higher $J_c$ of T-M1 compared to T-M2. That is also in agreement with observations in ZFC measurements shown in Figure 3. This tape presents indeed a very homogeneous magnetic flux profile and therefore only both images taken in zero-field are shown: in the decreasing (b) and increasing (c) field branch. Note that the Fe particles present in the surface of the core (Figure 3(b)-left) are well visible in MO images after field-cooling the sample, but do not have any influence in the superconducting behaviour of this tape, as expected.

The magnetic inhomogeneities observed in FC MO images of tape T-M2 in Figure 4 could be due to the presence of cavities or holes, or to the existence of poor superconducting links between regions within this sample. A further analysis by SEM of this tape discards the existence of cavities, and confirms the formation of aggregates of MgB$_2$ poorly connected with each other, which are observed in this part of the sample, as shown in Figure 6(b). The observed grain structure resembles the microstructure of some ex situ MgB$_2$ materials [24]. By contrast, other areas of this tape, with higher $J_c$ and good homogeneity according to MO images (as in the left area
in Figure 4), are well connected as it is seen in Figure 6(a). It must be remarked that the SEM images of tapes made with unmilled, Figure 6(c), and milled precursors with milling conditions M1, Figure 6(d), show well-connected superconducting grains.

![Figure 6. SEM images (secondary electrons detector) of the tape T-M2 in the left region (a), where $J_c$ is higher, and in the central area (b) where “bubble like shape” magnetic inhomogeneities are seen in Figure 4. Image (c) is of the tape with unmilled precursor and (d) of T-M1. The same magnification has been used in all images.](image)

5. Conclusions

We have analysed the effect of precursor ball milling on the properties of monocore Fe-sheathed MgB$_2$ wires and tapes prepared by the in situ powder-in-tube method by using drawing and rolling techniques. The effect of milling energies per unit mass in the range between $1.2 \times 10^6$ and $1 \times 10^8$ J kg$^{-1}$ (above the mechanical alloying limit) on the superconducting properties of these conductors were analysed.

Magneto-optical (MO) imaging analysis has been performed in order to investigate magnetic flux penetration in these samples, revealing local $J_c$ variations in these superconductors and allowing further understanding of the reasons of observed $J_c$ deterioration at low magnetic fields frequently observed with increase of the milling
energy of the precursor powders. It was found that microstructural characteristics change strongly with milling energy of the precursor and affect significantly the magnetic flux front propagation observed by MO imaging at zero-field-cooled and field-cooled conditions. MO imaging performed in field-cooled samples, following a specific magnetic field sequence, highlights the distortions in the magnetic pattern so that it has been found very helpful to reveal inhomogeneities in the superconductor. The MO images have been correlated with microstructural features observed with scanning electron microscopy (SEM), such as small cracks, sheath pores or weakly-linked aggregates of MgB$_2$ grains present in some of the analysed tapes.

The use of the precursor with milling energies above a certain limit produces microstructural defects during mechanical deformation, resulting in the formation of large transversal cracks, which indicates that using just drawing to deform the wire is not appropriate to fabricate conductors from high energy milled precursor powders. The tape fabricated from the precursor with the lowest milling energy shows good homogeneity but lower pinning at high fields and low temperatures than in tapes produced from higher-energy milled precursors.

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