






Effects of the oxygen source configuration on the superconducting properties of internally-oxidized internal-Sn Nb₃Sn wires

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Abstract

We successfully manufactured 12-filament rod-in-tube Nb₃Sn wires with oxide nanoparticles formed by the internal oxidation method. We employed Nb-7.5 wt%Ta-1 wt%Zr and Nb-7.5 wt%Ta-2 wt% Hf alloys along with oxygen sources (OSs) in two different configurations—in the core of Nb filaments (coreOS) and at the boundary between the filaments and the Cu tube (annularOS)—to assess the influence of the OS layout on the superconducting properties and grain size. The simultaneous presence of the OS and of Hf or Zr reduced the average Nb₃Sn grain size to around 50 nm, leading to an enhancement of the layer critical current density (J_c) up to 3000 A mm⁻² at 4.2 K and 16 T for the Hf-annularOS wire. Samples manufactured with an OS show a shift toward higher reduced magnetic fields of the position of the maximum in pinning-force density, this shift being more pronounced when SnO₂ is added in the annularOS configuration, and for the Hf-containing samples. This enhanced pinning at higher magnetic field is beneficial for high-field magnet applications. Moreover, we measured a very high upper critical field, reaching 29.3 T at 4.2 K in the Hf-annularOS samples.

Keywords: Nb₃Sn, internal oxidation, multi-filamentary wire

(Some figures may appear in colour only in the online journal)

1. Introduction

Nb₃Sn superconductor is currently the preferred solution for the conductor of the next generation of high field accelerator magnets. An example is the High Luminosity upgrade

of the Large Hadron Collider (HL-LHC) [1] at the European Organization for Nuclear Research (CERN). One of the objectives of this program is the installation in the interaction region of high-field large-aperture Nb₃Sn quadrupole magnets, with a peak field on the conductor of 12 T [2]. This goal required the development of Nb₃Sn wires with a non-Cu J_c (i.e., the critical current, I_c , divided by the wire's non-Cu area, including Nb₃Sn, residual bronze, and barriers) of 2450 A mm⁻² at 4.2 K and 12 T. CERN is undertaking a study for a new proton-proton collider known as the future circular collider (FCC-hh) [3], a ~100 km circumference ring with 16 T Nb₃Sn dipole magnets, and a resulting collision energy

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of 100 TeV. The need of reaching a dipole field of 16 T in a compact and economically viable magnet translates into a non-Cu J_c target for the Nb₃Sn wires of 1500 A mm⁻² at 4.2 K and 16 T [4]. For comparison, under the same conditions, the wires developed for the HL-LHC upgrade can exhibit a non-Cu J_c value as high as 1250 A mm⁻² at 4.2 K and 16 T [5, 6].

Progress in wire technology continuously improved the critical current density (layer J_c , i.e., the I_c divided by the Nb₃Sn area) until the early 2000s [7], when internal Sn multi-filamentary wires were optimized using the Rod-Restack-Process (RRP[®]) [8]. The RRP[®] is a rod-in-tube (RIT) technique that involves stacking hundreds of Nb filaments jacketed in Cu around a sacrificial Cu core that is subsequently replaced with a Sn rod to form a sub-element, which is later cut and restacked to create the multi-filamentary wire. A heat treatment at temperatures ranging from 640 °C to 700 °C is then needed to form the Nb₃Sn phase. In inclusion-free Nb₃Sn the grain boundaries are the main pinning centers; therefore, materials with finer grains have higher current densities. In highly optimized RIT wires, Nb₃Sn typically has grain sizes of ~100–150 nm [9]. Tailoring the heat treatment to promote the Sn diffusion and introducing proper amounts of Ti or Ta, it is possible to achieve non-Cu J_c values of 1200 A mm⁻² at 16 T and 4.2 K, whereas the fine tuning of the Cu to Nb local area ratio and the overall Nb content of the subelements [10] led to 1300 A mm⁻² (16 T, 4.2 K). These values are still 13%–20% lower than the FCC target [11]. Barzi *et al* [10] clearly concluded that the J_c of the RRP[®] wires has plateaued and that further efforts should concentrate on improving the inherent pinning of these wires. Similar considerations apply to the wires produced by the powder-in-tube (PIT) route [12], with the highest non-Cu J_c reported in the literature being close to 1300 A mm⁻² (16 T, 4.2 K) [13]. For the dependence of pinning force on grain size reported in the literature [14, 15], we can estimate that a reduction in the grain size down to 50–60 nm would effectively enhance the non-Cu J_c beyond the FCC specification.

Investigations on practical methods for inhibiting the grain growth in Nb₃Sn date back to the 1960s when the first evidence of Nb₃Sn grain refinement due to grain growth inhibition was observed by Benz [16]. A Nb-1%Zr tape was anodized and heat-treated to introduce oxygen in the Nb matrix. The resulting tape was then coated with Sn and heated above 930 °C to form a layer of Nb₃Sn at the interface. The presence of ZrO₂, formed by the internal oxidation of Zr in the alloy, inhibited the grain growth of Nb₃Sn. In composite multi-filamentary wires, adopting a similar approach is unfeasible because Nb filaments are embedded in a Cu or bronze matrix to lower the formation temperature of Nb₃Sn. Oxygen cannot readily diffuse into Nb from the outside at the end of the deformation process, and introducing oxygen into Nb beforehand would considerably increase its hardness and reduce deformability. The other possibility is to supply oxygen from a judiciously placed separate oxygen source (OS) from which oxygen can effectively diffuse through the Nb alloy after the deformation process (i.e. during heat treatment). Zeitlin *et al* [17] were the first to report an attempt to inhibit grain growth in a round-wire conductor by the addition of an OS. This was done in RIT type

wires based on Nb-1%Zr filaments whose solid Sn core was replaced by a mixture of Sn and SnO₂ powders. SnO₂ was supposed to decompose during the heat treatment and, after diffusion through the Cu and the Nb alloy, react with Zr (owing to its higher affinity for oxygen), forming ZrO₂ to inhibit grain growth. The reacted wire displayed no signs of grain refinement at the usual Nb₃Sn reaction heat treatment temperatures, presumably because of the Cu separating the Sn core from the filaments.

In 2014, Xu *et al* [18] demonstrated the feasibility of internal oxidation in a mono-filamentary tube-type wire, reporting Nb₃Sn with an average grain size of 43 nm. SnO₂ powder was added at the interface between the Nb-1%Zr tube and the internal Cu/Sn core. During heat treatment, the SnO₂ in contact with the Nb-1%Zr alloy decomposes, supplying oxygen to the alloy for the formation of ZrO₂ precipitates. The proposed mono-filamentary wire also exhibited a modified pinning mechanism, showing a shift in the maximum pinning force to higher reduced magnetic fields. At the University of Geneva, we were able to reproduce some of these results in mono-filamentary thin wires (diameter of 0.22 mm) with various Nb alloys (Nb-1 wt%Zr, Nb-7.5 wt%Ta-1 wt%Zr, Nb-7.5 wt%Ta-2 wt%Zr) and different OSs (SnO₂, CuO) that were centrally placed in the Nb alloy [19]. No pinning peak shift was observed, but samples fabricated with the Nb-7.5 wt%Ta-2 wt%Zr alloy exhibited an outstanding B_{c2} value of 29.2 T at 4.2 K [20]. Motowidlo *et al* [21] used a PIT method to fabricate multi-filamentary wires based on Nb-1 wt% Zr tubes filled with a mixture of Cu₅Sn₄ and Sn powders. A relatively large grain size gradient was observed, but the area equivalent mean diameter was as low as 30 nm at the interface with the Nb alloy. Transport measurements over a wide range of magnetic fields allowed the deconvolution of the pinning force density into grain boundary and point defect contributions, the latter being found to dominate with contributions from 65% to 76%, depending on composition and heat treatment. Balachandran *et al* [22] fabricated Nb₃Sn mono-filamentary wires based on a Nb alloy containing Hf, namely Nb-7.5 wt% Ta-1 wt% Hf. A rod of this alloy was placed in the core of the wire, surrounded by a mixture of Sn and Cu powder, and SnO₂ was added to the mixture to examine the effect of OS on different alloys. A Ta barrier was placed to hold the powder mix and prevent the contamination of the external Cu stabilizer. The use of a Hf-containing alloy without the intentional addition of OS reduced the Nb₃Sn average grain size from 107 nm (as measured on a reference Nb–Ta wire fabricated with the same architecture) to 68 nm. The wire fabricated with SnO₂ addition had a smaller grain size (55 nm) but exhibited a lower pinning force at the peak field. Banno *et al* [23] studied Nb₃Sn bulks made from Hf-containing Nb alloys, correlating the reduced Nb₃Sn grain size with the smaller grain size of the original Hf-containing Nb alloy. Grain refinement was observed in Nb₃Sn synthesized from an annealed alloy; however, the grain size of Nb₃Sn without intentional OS was reduced to 81 nm.

Xu *et al* [24] further applied the internal oxidation to powder-in-tube multi-filamentary Nb₃Sn wires based on ternary Nb alloys (Nb-7.5 wt% Ta-1 wt% Zr), with subelement numbers and wire lengths approaching those of practical

wires. To fabricate these ‘additional pinning center-powder in tube’ (APC-PIT) wires Cu-jacketed Nb-7.5 wt% Ta-1 wt% Zr tubes were filled with a mixture of Cu, SnO₂ and Sn powders. After the initial deformation, the resulting assemblies were cut and used as subelements in a Cu tube to prepare multi-filamentary wires. These multi-filamentary APC-PIT wires exhibited high B_{c2} owing to effective Ta doping [25], and high J_c owing to the effective ZrO₂ APC, some of them surpassing the FCC specification [26]. In other powder-in-tube multi-filamentary Nb₃Sn wires based on Nb-1 wt% Zr and SnO₂ powder as OS [27], grain refinement was not observed even though nanoscale precipitates were present and a shift of maximum pinning force density to higher magnetic field was measured. It was argued that the oxidation of Nb–Sn intermetallics deprives some of the Zr from oxygen needed for its oxidation.

In the framework of a collaboration between UNIGE and CERN, we present here the results of our investigation on internal Sn wires with an internal OS to enhance the superconducting properties of Nb₃Sn. This study explores methods to add an internal OS to a simplified RIT wire, which imitates a subelement of RIT wires with a reduced number of Nb-alloy filaments. The absence of powder in the design of current RIT wires necessitates preliminary studies to evaluate the effects of powder OS in different layouts on wire deformability/fabrication and superconducting properties. We have developed a versatile design that enables us to investigate the properties of different Nb-alloys (Nb-7.5 wt% Ta-1 wt% Zr; Nb-7.5 wt% Ta-2 wt% Hf) combined with two possible OS configurations (*annularOS* and *coreOS*) in a 12-filament arrangement surrounding a Sn core. This fast turnaround R&D activity allowed us to investigate the effects on the superconducting properties of the position of the OS relative to the Nb₃Sn formation front, as detailed below.

2. Experimental methods and details

2.1. Wire fabrication

The wires presented in this work were based on two different Nb-7.5 wt%Ta alloys, distinguished by a third element:

- (1) Hf in the Nb alloy: Nb-7.5 wt%Ta-2 wt%Hf
- (2) Zr in the Nb alloy: Nb-7.5 wt%Ta-1 wt%Zr.

The weight percentage of Hf was selected such as to have the same atomic fraction of Hf in the Nb–Ta–Hf alloy as of Zr in the Nb–Ta–Zr alloy, namely 1.1 atomic %.

For each of these alloys, we prepared a reference wire without an OS and two wires with different OS layouts, as illustrated in figure 1 and described here:

- (a) *noOS*: without OS as reference sample for the series (figure 1(a))
- (b) *coreOS*: SnO₂ was placed at the core of each Nb alloy filament (figure 1(b))
- (c) *annularOS*: SnO₂ placed around each Nb-alloy filament (figure 1(c)).

We additionally manufactured a wire using a Nb-7.5 wt%Ta alloy, without OS, used as a reference sample for comparison (NbTa-*noOS*). Table 1 provides a summary of the manufactured wires with the alloy used for the filaments and OS layout.

To prepare the filaments of the *coreOS* wires, a central hole was drilled in the Nb alloy and filled with SnO₂ powder (Inframat Advanced Materials, 80 nm nominal particle size) at tap density. The hole was sized to accommodate enough OS to supply the oxygen necessary for oxidizing all the Hf or Zr atoms in the Nb alloy, assuming a 50% packing density of the oxide powder.

For the *annularOS* technique, we coated the Nb alloy filaments with a thin layer of SnO₂ by dip coating [28]. Colloidal solutions of the SnO₂ (30 wt%) in ethanol were prepared, and included polyvinyl butyrate (PVB, 2 wt%) to reduce the formation of cracks in the coated layer during the evaporation of the ethanol. The Nb alloy rods were dipped into the suspension and air-dried for 10 min. This process was repeated until the deposited SnO₂ (evaluated by weighing the dried rod) was sufficient to provide the amount of oxygen required to oxidize all Hf or Zr atoms in the Nb alloy.

The Cu/Nb filaments were assembled by inserting a 4.9 mm Nb alloy rod (with or without an OS) into a Cu tube (outer diameter, OD = 6.30 mm; inner diameter, ID = 5.30 mm). The assembly was cold-deformed by swaging and wire drawing to a diameter of 1.12 mm. For reasons related to the co-deformation of the Cu/Nb assembly, we used an excess of Cu relative to the desired Cu-to-filament area ratio of 0.2. After the deformation process, electropolishing [29] was employed to remove excess Cu from the Cu/Nb filaments.

The Cu/Sn rod was directly assembled with the intended Cu-to-filament area ratio of 0.2.

Twelve Cu/Nb filaments and a central Cu-clad Sn rod were inserted inside a Ta tube, which acts as a barrier to preserve the purity of the external Cu stabilizer. The surfaces of all the filaments and tubes were mechanically and chemically cleaned before assembly. The assemblies were then swaged and drawn to achieve a final diameter of 0.70 mm.

After cleaning, straight samples (16 cm in length) with hammered ends (to minimize Sn leakage) were inserted into quartz tubes and heat treated under vacuum at 650 °C for 200 h, with an intermediate plateau at 550 °C for 100 h. This plateau at 550 °C is thought to facilitate oxygen diffusion into the Nb alloy before the formation of Nb₃Sn, as also used in [22].

2.2. Characterization techniques

After deformation, we cut short wire pieces to examine the cross-section of the deformed wire and assess the filament homogeneity using scanning electron microscopy (SEM) images taken with a JEOL JSM-7600F field emission scanning electron microscope (FESEM). For each filament in a wire cross-section, we measured the average of four diameters along four different directions. Taking into account all 12 filaments, we calculated the mean average filament size for each wire and its standard deviation. Moreover, we measured the

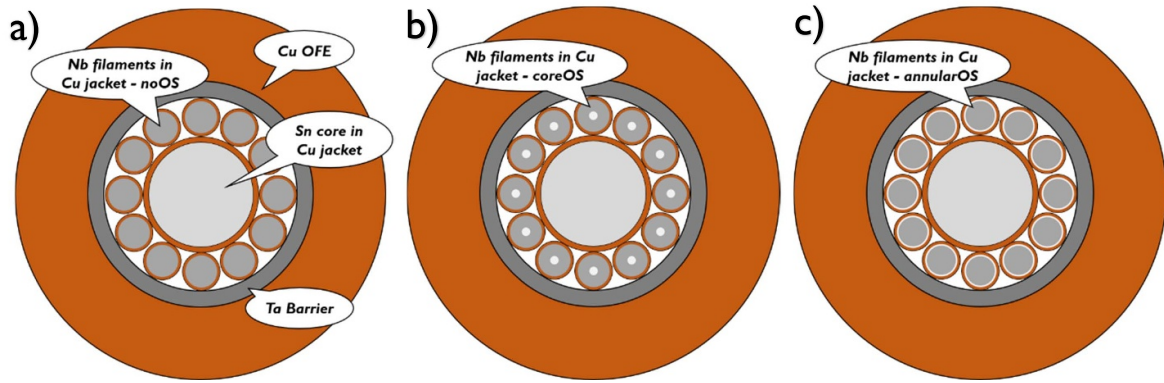


Figure 1. Rod-in-tube design for the wires reported in the paper and comparison between SnO₂ layouts: (a) no addition of SnO₂ (*noOS*), (b) SnO₂ added in the Nb filament core (*coreOS*), or (c) SnO₂ at the interface between Nb filaments and Cu jacket (*annularOS*).

Table 1. List of wire types and their filament alloy and oxygen source configuration. All percentages are by weight.

Wire type ID	Alloy in the Cu/Nb filaments (wt%)	Oxygen source material	Oxygen source position
NbTa- <i>noOS</i>	Nb-7.5%Ta	No oxide	N.A.
Hf- <i>noOS</i>	Nb-7.5%Ta-2%Hf	No oxide	N.A.
Hf- <i>coreOS</i>	Nb-7.5%Ta-2%Hf	SnO ₂	Core
Hf- <i>annularOS</i>	Nb-7.5%Ta-2%Hf	SnO ₂	Annular
Zr- <i>noOS</i>	Nb-7.5%Ta-1%Zr	No oxide	N.A.
Zr- <i>coreOS</i>	Nb-7.5%Ta-1%Zr	SnO ₂	Core
Zr- <i>annularOS</i>	Nb-7.5%Ta-1%Zr	SnO ₂	Annular

areas and elemental area ratios of Cu, Nb, and Sn enclosed by the barrier (excluding external stabilizer Cu and barrier).

Following the heat-treatment, we examined fractured cross-sections of the wires to determine the average grain size following the line intercept method. As also observed in our previous work [20], wires containing an OS exhibited elongated Nb₃Sn grains, so we evaluated the grain size in both the long- and short-axis directions. We determined the intercept distances of two lines for each axis direction. This was repeated on multiple images (from 5 to 10) for each wire to calculate the mean intercept distance (as an arithmetic mean) and its standard deviation.

Polished cross-sections were examined using the FESEM to determine the thickness and the area of the fine Nb₃Sn grain region. We calculated the thickness as the arithmetic mean over multiple values of the thickness of the fine-grained Nb₃Sn area. For each reacted filament we measured the length of four diameters along four different directions to calculate the outer (D) and inner diameter (d) of the fine-grained area. The difference divided by two was used to calculate the average layer thickness. The area of fine Nb₃Sn grains was measured on the wire samples subjected to transport critical current (I_c) measurement.

I_c measurements were performed at 4.2 K in magnetic fields of up to 19 T at the University of Geneva. The samples were 50 mm long, with two sets of voltage taps placed at distances of ~ 20 and ~ 10 mm. The criterion for the critical current determination was $0.1 \mu\text{V cm}^{-1}$. The layer J_c was calculated by dividing I_c by the transverse area of fine-grained Nb₃Sn. The determination of I_c was only possible down to 10–12 T, due to the presence of electromagnetic and thermal instabilities at

lower fields. As a complement, the layer J_c was extracted using equation (1) from the $M(H)$ loops acquired in a temperature range from 4.2 to 16 K and in magnetic fields up to 7 T using a VSM SQUID. The formula is an application of Bean's critical state model [30] to a hollow cylinder [31, 32] with its axis perpendicular to the magnetic field [33]:

$$J_{c,\text{layer}} = \frac{30\pi}{4} \times \frac{1}{1 - \left(\frac{d}{D}\right)^3} \times \frac{\Delta M}{D}. \quad (1)$$

The opening (ΔM) of the $M(H)$ loop in the magnetic field of interest was measured in emu cm^{-3} . D and d represent the outer and inner diameters (cm) of the fine-grained area of the superconductor in the shape of a hollow cylinder, respectively. The values of $J_c(B, T)$ were used to calculate the pinning force ($F_p = J_c \times B$).

We determined the B_{c2} as a function of temperature using low-current four-point transport measurements in magnetic fields up to 35 T at the Laboratoire National des Champs Magnétiques Intenses (LNCMI) in Grenoble, France, on a specially designed four-sample probe [34]. The samples were probed with a current between 1 and 5 mA, equivalent to a current density in the superconductor in the range of 0.1 – 0.5 A mm^{-2} . The upper critical field was extracted from the $R(B)$ curves using different criteria (99%, 90%, 50%, 10%, and 1%). For most samples, the outer Cu was chemically removed to expose the Ta barrier, which increased the electric resistance of the samples and enhanced the signal-to-noise ratio. Several wire samples were measured both with and without external Cu, demonstrating a negligible effect of precompression on B_{c2} on the wires manufactured for this work.

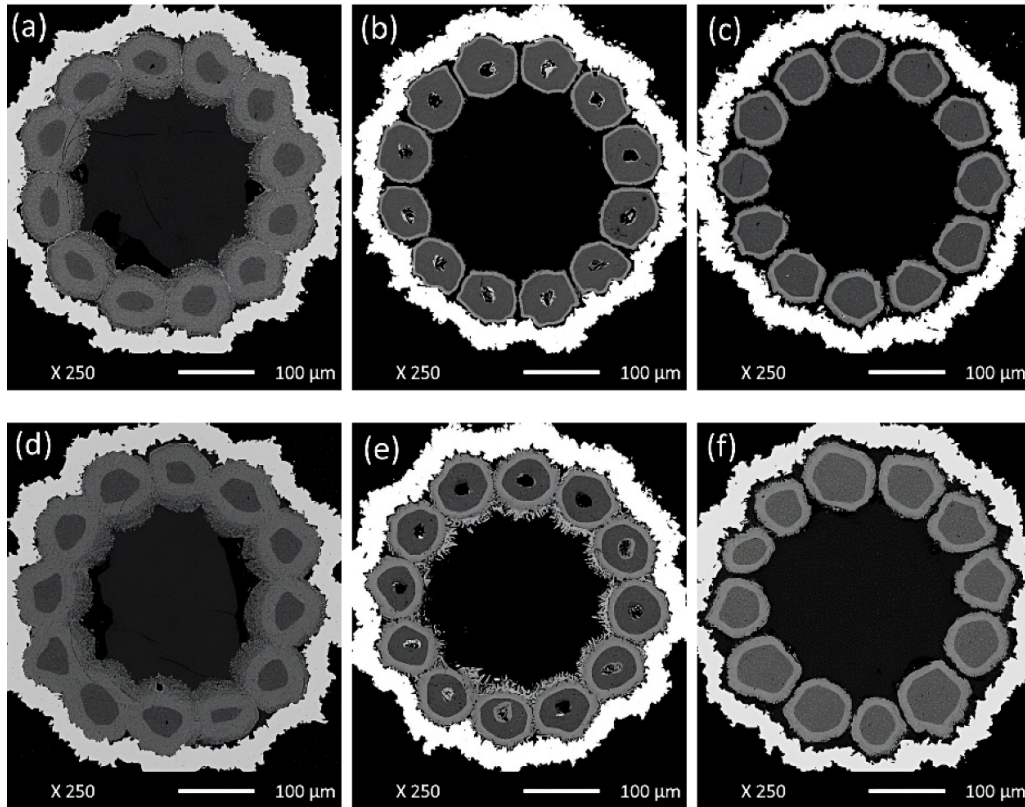


Figure 2. Back-scattered electron images of polished transverse cross-sections of wire samples after heat treatment: Hf-*noOS* (a), Hf-*coreOS* (b), Hf-*annularOS* (c), Zr-*noOS* (d), Zr-*coreOS* (e), Zr-*annularOS* (f).

Table 2. Summary of wire properties (elemental ratio in the cross-section of unreacted wires, filament size) and microstructure properties of reacted samples (fine-grained layer thickness and Nb₃Sn grain size). Elemental ratio is expressed relative to the total area inside the barrier (TOT), excluding barrier and external Cu. Standard deviations are given as a measure of the dispersion around the mean value.

Wire ID	Nb/Sn	Sn/Cu	Sn/TOT	Nb/TOT	Cu/TOT	Average filament size (μm)	Fine grains layer thickness (μm)	Long axis grain size (nm)	Short axis grain size (nm)
NbTa- <i>noOS</i>	1.22	1.45	0.34	0.42	0.23	76 \pm 8	18 \pm 3	119 \pm 8	113 \pm 9
Hf- <i>noOS</i>	1.55	1.53	0.31	0.49	0.20	89 \pm 9	23 \pm 2	120 \pm 9	106 \pm 8
Hf- <i>coreOS</i>	2.06	0.82	0.23	0.48	0.29	73 \pm 3	04 \pm 1	054 \pm 6	040 \pm 6
Hf- <i>annularOS</i>	1.44	1.00	0.29	0.42	0.29	67 \pm 5	09 \pm 2	060 \pm 3	038 \pm 5
Zr- <i>noOS</i>	1.68	1.63	0.30	0.51	0.19	90 \pm 8	23 \pm 3	112 \pm 8	097 \pm 5
Zr- <i>coreOS</i>	1.57	1.52	0.31	0.49	0.20	81 \pm 3	13 \pm 2	0072 \pm 12	052 \pm 9
Zr- <i>annularOS</i>	1.42	1.30	0.31	0.45	0.24	086 \pm 11	10 \pm 2	078 \pm 6	057 \pm 5

3. Results and discussion

Figure 2 shows polished transverse cross-sections after the reaction heat treatment of wire samples containing Hf (from 2.a to 2.c), and Zr (from 2.d to 2.f). Further information regarding our wires, including elemental area fraction in unreacted wires and fine grain layer thickness of the manufactured wires, is reported in table 2. The thickness of the fine-grained area (fine grains layer thickness in table 2, visible as the light grey annular region at the periphery of the reacted filaments in figure 2) varies from wire type to wire type, with a drastic reduction from $\sim 20 \mu\text{m}$ to less than $10 \mu\text{m}$ when an OS is present. Considering a typical RIT wire developed for HL-LHC, whose final Nb filament size in a subelement is

approximately $1 \mu\text{m}$ and layer thickness is about $10 \mu\text{m}$ [11], we can argue that this important reduction of the diffusion kinetics in wires with an OS should not be a concern. However, further investigation in real scale multifilamentary wire will be necessary to make sure that the subelement can be fully reacted on a reasonable timescale.

Figure 3 shows representative images of the Nb₃Sn grains for each wire. As expected, the *noOS* wires prepared with Hf or Zr (figures 3(a) and (d)) exhibit large grains above 100 nm on both axes. Contrary to the findings of Balachandran *et al* [22] and Banno *et al* [23], our samples do not show grain refinement induced solely by the presence of Hf in the Nb alloy, as the grain size is consistent with that of commercially available Nb₃Sn wires and our own NbTa-*noOS* samples. The

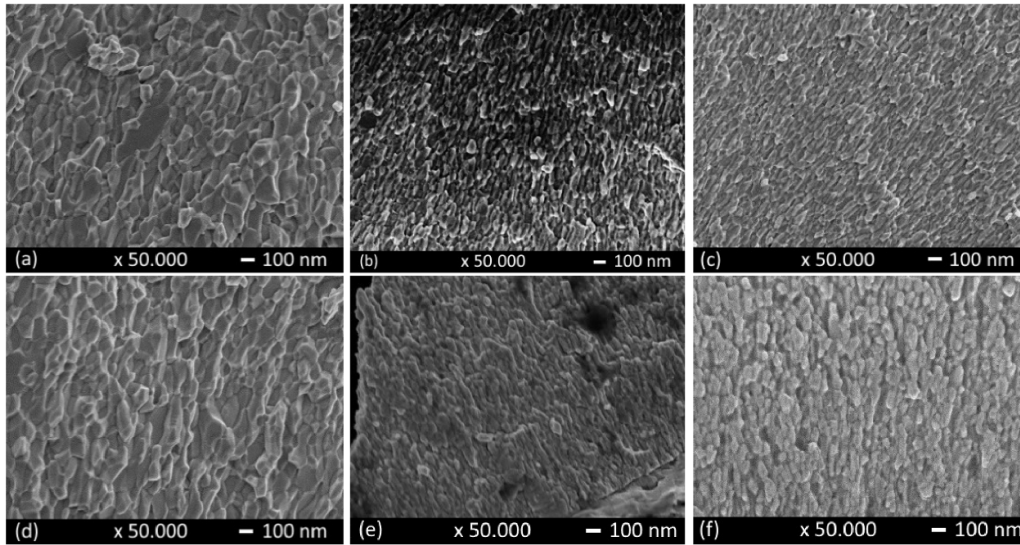


Figure 3. SEM images of fractured cross-section of Hf and Zr-containing wires: HfnoOS (a), Hf-coreOS (b), Hf-annularOS (c), Zr-noOS (d), Zr-coreOS (e), Zr-annularOS (f).

Hf-coreOS (figure 3(b)) and Hf-annularOS (figure 3(c)) wires exhibit a more elongated shape of the Nb₃Sn grain with respect to Hf-noOS wire as well as a pronounced reduction of the grain size. The mean size in the short axis direction was reduced to less than 40 nm for both of these wire types. Grain refinement was also observed in the Zr-containing samples when OS was added, with an elongated shape. However, for the Zr-containing Nb₃Sn, the effect of grain refinement induced by OS was less pronounced than that of the Hf-containing samples. Along the short axis, grains are refined down to 52 and 57 nm for Zr-coreOS and Zr-annularOS respectively. For both Hf and Zr samples, there is no significant difference in terms of grain refinement between the core and the annular placement of the OS.

At higher magnifications, brighter dots were observed on the exposed grains, as shown in figure 4. Based on previous observations and analyses [26, 35, 36], these dots are likely oxide nanoprecipitates (HfO₂ or ZrO₂, depending on the third element added to the Nb alloy). Currently we are not able to determine the composition or exact size of these particles. Based on these images, it would appear that these nanoparticles are mostly present at the grain boundaries, but it should be noted that the SEM images of the fractured samples reveal only oxide particles at the grain boundaries, leaving hidden oxide nanoparticles that may be present inside the grains. Transmission electron microscopy and atomic probe tomography performed by Xu *et al* [26, 37] have shown that, in average, there are more than ten oxide nanoparticles in each Nb₃Sn grain.

Samples containing an OS exhibited a significant improvement in layer J_c compared to noOS wires, as shown in figure 5. To underline the magnitude of the J_c enhancement, we added to the graph the layer J_c that would lead to a non-Cu J_c of 1500 A mm⁻² (at 16 T, 4.2 K, the FCC target) in a wire with the same layer fraction of 60% of the non-Cu area as the high performance wires developed in the frame of the

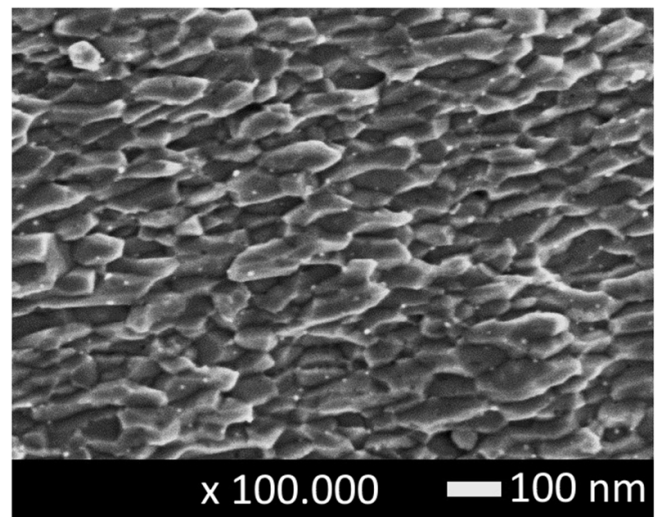


Figure 4. High magnification image of a fractured Hf-coreOS sample, showing oxide precipitates visible as bright spots.

HL-LHC project [2, 22]. This layer J_c amounts to an effective FCC layer target of 2500 A mm⁻² at 16 T and 4.2 K. All the samples fabricated with an internal OS exhibit layer J_c values above this target. In particular, the Hf-annularOS wire exhibits a layer J_c of 3000 A mm⁻² at 16 T and 4.2 K, while the samples containing Zr reach 2700 A mm⁻². While these J_c values are quite high compared with those in current state of the art Nb₃Sn wires without internally oxidized nanoparticles they are still well below the values exceeding 4000 A mm⁻² that were measured by Xu *et al* [38] in PIT-APC Nb₃Sn wires based on Nb-Ta-Hf and Nb-Ta-Zr of the same nominal compositions as the alloys of our wires. The grain size distributions appearing comparable we can attribute this difference in J_c either to the distribution of the number concentration of the oxide nanoparticles and/or to distribution of more fundamental

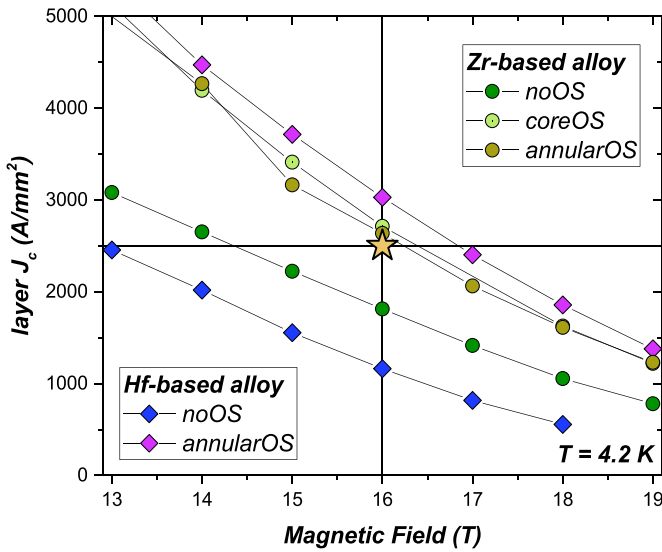


Figure 5. Layer J_c of the wire samples containing Hf or Zr, with and without OS. The star represents the FCC target as layer J_c in a wire having the same layer to non-Cu volume ratio as a HL-LHC RRP[®] wire.

properties like B_{c2} . We lack currently the technical means to characterize these distributions but we think that most of this difference comes from the gradient of the Sn content in the Nb_3Sn layer, which has a marked influence on B_{c2} distribution. We expect that RIT wires with internally oxidized nanoparticles and finer filaments heat treated to obtain lower composition gradients across the layer would also exceed the 4000 A mm^{-2} at 16 T and 4.2 K. In this context it is important to note that Xu *et al* [38] have also observed important differences between the J_c -s of wires based on the same alloy and same oxygen supply but different Nb/Sn ratios. As a further proof of the importance of other design parameters on the properties of the reacted layer we observe that because of the very thin Nb_3Sn layer (as shown in figure 2(b) and reported in table 2), the *coreOS* wire samples containing Hf did not have a sufficiently continuous layer of superconductor to allow current transport measurements.

The enhancement of J_c exhibited by the internally oxidized wires is not only due to grain refinement but also to a significant enhancement of B_{c2} . Figure 6 shows the ranges of B_{c2} values at different criteria as measured on multiple samples of each wire type. Experimental data show that B_{c2} (4.2 K) calculated using the 99% criterion increases to 29.2 and 29.3 T for Zr-*annularOS* and Hf-*annularOS* samples, respectively, as shown in figure 7. These values exceed by more than 1 T the B_{c2} value of 28 T for a sample fabricated with the same procedure and using Nb-7.5 wt%Ta alloy (NbTa-*noOS*). These results are in agreement with our previous measurements of mono-filamentary wires [20]. Considering the $B_{c2}(99\%)$, it can be seen that for both Hf- and Zr-containing samples, the introduction of OS enhances B_{c2} progressively from *coreOS* to *annularOS*. Zr-*noOS* did not show any B_{c2} enhancement with respect to the NbTa-*noOS* wire used as a reference, whereas the presence of Hf enhanced B_{c2} already without an OS. All our Zr-containing samples show a similar transition width,

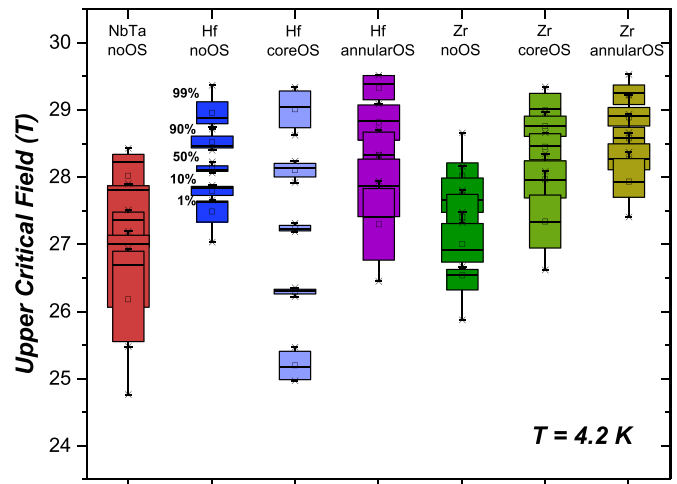


Figure 6. B_{c2} (4.2 K) values of several samples of each wire calculated with different criteria (from top to bottom: 99%, 90%, 50%, 10% and 1%). Comparison with wire NbTa-*noOS*, fabricated with the same procedure using Nb-7.5 wt%Ta, is given.

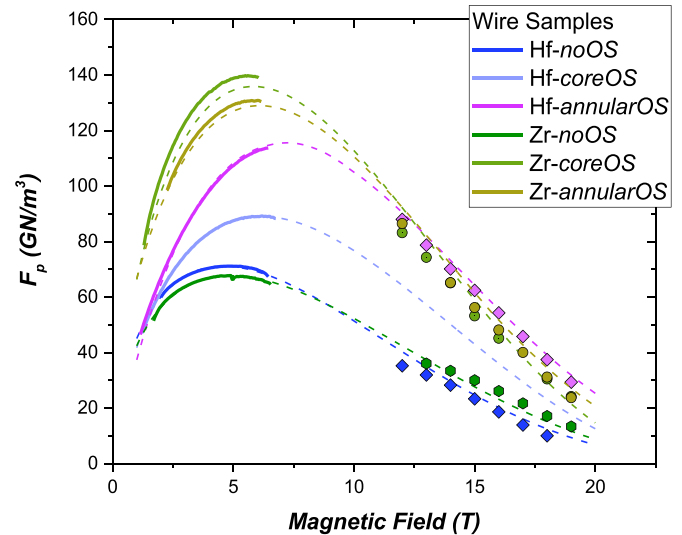


Figure 7. Pinning force as a function of magnetic field for Hf and Zr-containing samples, calculated from magnetic and transport layer J_c .

according to the difference between 99% and 1% criteria (i.e. $B_{c2}(99\%) - B_{c2}(1\%)$) reported in table 3, with no significant changes with introduction of OS. The Hf-containing samples exhibited higher variability with a wider transition width when OS was added.

These results confirm the enhancement of B_{c2} in the presence of Zr and OS reported in our previous work [20], and in the presence of Hf, as reported by Xu *et al* [18]. Thus far, a possible explanation given by Xu *et al* [37] argued that the presence of ZrO_2 (or HfO_2) leads to a higher Sn content in Nb_3Sn owing to the suppressed reaction rate at the Nb_3Sn/Nb alloy interface. A higher Sn concentration in Nb_3Sn , which is closer to stoichiometry, enhances B_{c2} . The reduced reaction layer observed in the cross-sections (figure 2) of the wires with an OS supports the explanation proposed by Xu *et al* [37].

Table 3. Average B_{c2} (99 and 1%) and F_p peak positions at 4.2 K.

Wire ID	Average B_{c2} (99%) (T)	Average B_{c2} (1%) (T)	$B/B_{c2}(1\%)$ at $F_p(\text{max})$
NbTa- <i>noOS</i>	28.0	26.1	0.19
Hf- <i>noOS</i>	28.9	27.6	0.20
Hf- <i>coreOS</i>	29.0	25.1	0.24
Hf- <i>annularOS</i>	29.3	27.3	0.30
Zr- <i>noOS</i>	28.0	26.5	0.20
Zr- <i>coreOS</i>	29.0	27.3	0.22
Zr- <i>annularOS</i>	29.2	27.9	0.24

However, a more detailed investigation on the possible role of precipitates will be needed to understand their role in the B_{c2} enhancement, and if the same B_{c2} values will be observed in a real RIT wire with internal oxidation.

When an OS is added, a shift in the F_p peak ($F_p = J_c \times B$) to a higher absolute magnetic field is observed, as shown in figure 7, which shows the F_p curves calculated from both magnetic and transport measurements. The fits (dashed lines) were obtained using the B_{c2} (1%) in table 3 as a fixed parameter. This shift in the F_p peak is more substantial in the samples containing Hf, with the Hf-*annularOS* sample showing a peak at 7.3 T. The introduction of the OS also leads to an enhancement in the maximum value of F_p . At the peak, the Zr-containing samples with OS exhibits higher values of F_p of 140 and 130 GN m⁻³ for *coreOS* and *annularOS* respectively. Without OS, the Hf-containing sample exhibits a maximum F_p of 70 GN m⁻³, which is progressively enhanced by the introduction of OS to 90 GN m⁻³ in the *coreOS* sample and to 135 GN m⁻³ in the *annularOS* sample. At a higher magnetic field, the Hf-*annularOS* sample takes over and exhibits the highest F_p at 16 T.

The shift of F_p peak is driven by the enhancement of B_{c2} as shown in figure 6, and by a modification of the pinning mechanism induced by the presence of HfO₂ and ZrO₂ precipitates. This can be observed by analyzing the normalized F_p as a function of reduced field ($b = B/B_{c2}(1\%)$) reported in figure 8.

Figure 8(a) shows the *normalized* F_p curves of the Hf-containing samples and figure 8(b) those of the Zr-containing samples, all calculated from the inductive $J_c(B)$ curves measured between 4.2 and 14 K. The magnetic field values were scaled to $B_{c2}(1\%)$ (as obtained from the $R(B)$ measurements) because this criterion gives a better scaling of the F_p , as compared with the 10% criterion. Wire samples without an OS display a peak of the normalized F_p at 0.2, as expected in the case of grain boundary pinning [39]. The introduction of the OS induces a shift in the F_p peak, which is more pronounced for the *annularOS* wires in both the Hf- and Zr-containing samples. The most significant shift of the peak is seen in the Hf-*annularOS* sample, with the peak occurring at 0.30 of the reduced field, which is attributable to an additional contribution to the pinning, from a different type of defect than the grain boundaries. An F_p peak at $b = 0.33$ can be associated with pinning by normal point defects [39]. The oxide nanoparticles are the obvious candidates for such pinning, and this is the chosen explanation given by Motowidlo *et al* [21],

Buehler *et al* [27] and Ortino *et al* [40]. In view of their results on samples reacted at low temperatures, with a particularly low average grain size of 30 nm and for which F_p maximized at $b = \sim 0.4$, Xu [9] concluded that instead of point pinning, individual pinning becomes important as the grain sizes approach the flux line spacing. If this were the actual additional pinning mechanism, a further reduction of the grain sizes would push the F_p peak towards the theoretical $b = 0.5$, a very interesting situation for high field applications. However, in more recent publications [24, 26], the same authors appear to favor the point defect pinning as an explanation for the shift of the F_p peak. A priori nothing excludes the possibility of three pinning contributions, but only further research will clear this issue. Moreover, based on the analysis of the change in the height of the F_p peak when passing from binary to Ta-doped Nb₃Sn, Ortino *et al* [40] cast doubt on the direct summation of grain boundary and point defect pinning.

In the absence of a clear understanding of the different pinning contribution, we can only enumerate and discuss the properties that may cause the difference observed between the positions of the F_p peaks of our Hf- and Zr-containing samples: the size distribution of the oxide precipitates, the spatial distribution of the oxide precipitates, the size distribution of the Nb₃Sn grains or a combination of some of these properties. Theoretical calculations show that the optimal size for flux pinning on insulating inclusions, like the oxide precipitates in these wires, is approximately twice the coherence length [41] of the superconductor. According to Xu *et al* [42], the element added to the Nb alloy has a significant influence on the oxide precipitate size (under the same heat treatment conditions), reporting that HfO₂ precipitates are smaller in size with respect to ZrO₂.

The grain refinement in Nb₃Sn with internally oxidized nanoparticles is due to the Zener drag effect [43], according to which the mobility of the grain boundaries, and thus the growth of the Nb₃Sn grain size, is reduced by the presence of small-sized precipitates, the resulting grain size being smaller for a low size and a high volume fraction of these precipitates. A high density of these precipitates can thus not only enhance the contribution to the pinning from the grain boundaries (finer grains imply a higher grain-boundary density) but also lead to a more important contribution to the vortex pinning from the nano-precipitates themselves, which results in a shift towards $b = 0.33$ of the F_p peak. Xu *et al* [42] show that the internal oxidation of Hf-containing samples produces larger effects on J_c compared to Zr, as HfO₂ ends up with a higher number concentration than ZrO₂ at the same solute atomic concentration in the starting Nb alloy, due to the smaller precipitate size of the HfO₂. The smaller size of the HfO₂ precipitates in comparison with ZrO₂ leads to a more effective Zener drag whereas the related higher number concentration tends to enhance the point defect pinning contribution. Comparing wires of the same atomic fraction of Hf and Zr (which is the case for Xu *et al* [42] and for us) and assuming complete internal oxidation of these species, the above considerations are experimentally supported by the smaller Nb₃Sn grains observed in Hf-containing samples compared to Zr-containing samples: Xu *et al* [42] observed Nb₃Sn grains of 53 nm and 64 nm in the

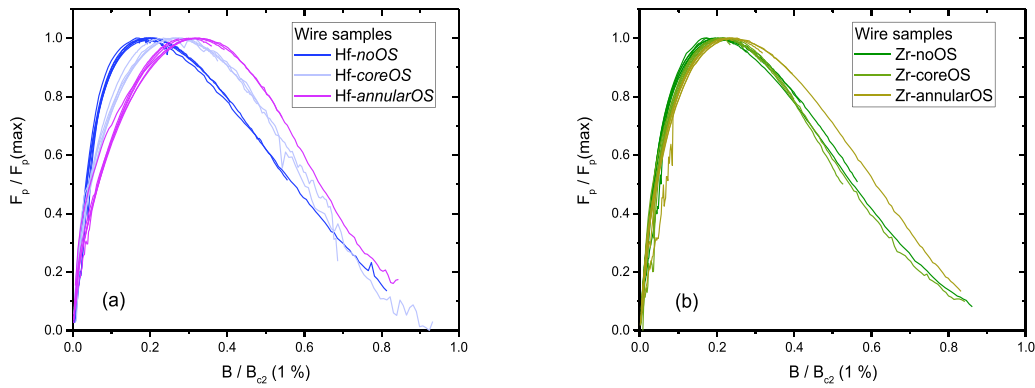


Figure 8. Normalized pinning force as a function of reduced field for Hf (a) and Zr-containing (b) samples. $B_{c2}(1\%)$ from $R(B)$ measurements is used to normalize the magnetic field.

outer boundaries for Hf and Zr samples (reacted at 675 °C), respectively, just as we observed smaller Nb_3Sn grain sizes in the Hf-containing samples with respect to the Zr-containing ones. However, in the absence of full characterizations of the size and spatial distributions we run the risk of looking at correlations rather than causalities. The F_p peak shift in our samples follows the same trend of higher values for HfO_2 but the peak values of F_p are less consistent. Unfortunately, other publications reporting measured dimensions of oxide precipitates do not allow fair comparisons as they are made on very different samples (manufacturing techniques, starting alloys and OS). They appear nevertheless to confirm a lower size for the HfO_2 precipitates. Based on atomic probe tomography, Tarantini *et al* [36] reported HfO_2 precipitate sizes of approximately 3 nm and Nb_3Sn grains below 70 nm (between 55 and 68 nm) in a wire manufactured by Balachandran *et al* [22] and reacted at 670 °C. Ortino *et al* [40] measured by transmission electron microscopy ZrO_2 precipitates of 4.6 nm and Nb_3Sn grain sizes of 70 nm in APC-PIT reacted at 675 °C. This comparison based on literature data and our own results in this paper is unfortunately not sufficient to fully understand how the size and distribution of oxides precipitate affect the grain refinement. Moreover, the variation of the size distributions of the Nb_3Sn grain across the reacted layer have rarely been systematically studied. One of these reports [21], shows not only significant grain size gradients but also broad size distributions. Investigations looking at the distributions of the precipitate size and precipitate position, and at the grain size vs position would be necessary to fully understand the influence of the precipitate particle size and number distributions on the grain refinement and ultimately on flux line pinning.

The present investigation shows that the placement of the OS influences the superconducting properties, but has a minimal impact on the grain refinement, as both approaches (core and annular) reduce the Nb_3Sn grain size to similar values for a given type of Nb alloy. The annular approach for introducing the OS yields better results in terms of B_{c2} enhancement and shift of the F_p peak position compared to the core approach. The *coreOS* samples have nevertheless significantly better properties than the wires manufactured without an OS,

underlining that an OS is essential for enhancing the high-field properties, whereas its placement enables optimization of the F_p and B_{c2} . The reasons for the different behaviors of the OS layouts are not yet clear but our future investigations will be aimed at determining whether they are related to different size and number distributions of the oxide nanoparticles resulting from different oxygen diffusion gradients.

4. Conclusions

We reported the successful implementation of the internal oxidation process on simple multi-filamentary wires manufactured through the RIT approach. Hf or Zr was added to the Nb alloy while SnO_2 was used as OS in two different configurations: core and annular. The introduction of the OS resulted in a refinement of the Nb_3Sn grains, which have a slightly elongated shape. In the wires with Hf and OS, the grain size was reduced to values below 40 nm on the short axis and below 60 nm on the long axis. Only a minor difference in grain size was observed between the *coreOS* and the *annularOS* layouts. In *Hf-annularOS* wires, the grain refinement produced an enhancement of the layer J_c up to 3000 A mm⁻² at 16 T and 4.2 K. This value exceeds the layer J_c of 2500 A mm⁻² necessary to achieve a non-Cu J_c of 1500 A mm⁻² (at 16 T, 4.2 K, the FCC target) in a wire having the same layer fraction as the high-performance wires developed for the HL-LHC project. The Hf or Zr oxide precipitates not only inhibit the grain growth during the heat treatment but also appear to act as effective pinning centers, causing a shift in the pinning force peak to higher reduced fields. This shift is more pronounced when SnO_2 is added in the *annularOS* configuration, and for the Hf-containing samples. Indeed, the *Hf-annularOS* sample shows an F_p peak at a field value corresponding to 0.3 B_{c2} . The introduction of OS resulted in $B_{c2}(99\%, 4.2 K)$ values of 29.2 and 29.3 T for Zr and *Hf-annularOS* samples respectively. These values are approximately 1.3 T higher than those of a wire manufactured with a Nb–Ta alloy, which was used as a reference. The mechanism behind the enhancement of B_{c2} and the different impact of the *coreOS* and *annularOS*

approaches on the F_p -peak shift are not fully understood and deserve dedicated investigations. On a general perspective, this work presents a promising approach to introduce a powder OS in a simplified RIT wire. This is a first step towards the introduction of an OS in the layout of the subelement of practical RIT wires, which presents difficulties related to the fine size of the filaments in the final wires and to the need for a uniformly distributed OS that does not affect negatively the fabrication of the wire. In this study, we examined the annular OS and the core OS configurations, given their previous use in other works [18, 20]. However, alternative OS configurations will probably be more suitable for RIT wires. In our forthcoming research, we will explore various methods for incorporating an OS into full-scale RIT subelements and examine their effect on deformability and superconducting properties. This is currently the major technological challenge for integrating internal oxidation into RIT wires. Given that a winning solution has not been yet identified, it is too early to evaluate how significantly this will increase the conductor cost but it is fairly clear that the additional manufacturing steps could bring it closer to the cost of the PIT-type Nb₃Sn superconductors. The fabrication of PIT-APC Nb₃Sn wires is uniquely suitable for the introduction of an OS but we think that continued research and development on both RIT and PIT-APC Nb₃Sn wires is the best way to ensure the availability of suitable conductors for future large scale projects like FCC.

Data availability statement

The data that support the findings of this study are openly available at the following URL/DOI: Data will be available upon request.

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