QUANTITATIVE MICROSTRUCTURAL ANALYSIS AND PIEZOELECTRICITY OF HIGHLY DENSE, SUBMICRON-STRUCTURED NaNbO₃ CERAMICS FROM MECHANICALLY ACTIVATED PRECURSORS

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Abstract

We study in this work the processing of NaNbO₃ ceramics prepared in a single thermal treatment of highly reactive precursors obtained by mechanical activation of different reagents, aiming to determine optimum conditions for piezoelectric ceramics production. Pressure-less sintering at 1200°C leads to dense ceramics (<5% porosity) with poor mechanical stability, unsuitable for practical uses. Dense hot pressed ceramics were also obtained at lower temperatures (900-1100°C), all of them in the submicron range of average grain sizes (< 400 nm). Their microstructure was quantitatively characterised and their elastic and electromechanical properties determined by an automatic iterative method from impedance measurements at resonance. A noticeable ensemble of piezoelectric and elastic properties (d₃₃=38 pC.N⁻¹ and Nₚ=3252 kHz.mm⁻¹) was measured for hot pressed ceramics, from precursors obtained by a combined route of wet-chemistry and mechanical activation, with a microstructure characterized by 0.4% residual porosity and a bimodal lognormal distribution of grain size.

Keywords: Quantitative Microstructure Analysis, Mechanical activation, Piezoelectricity, Niobates.

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INTRODUCTION

There is a recent need for finding environmentally friendly lead free piezoceramics with properties comparable with the ones of the commercial piezoceramics, based mainly in lead compounds. In this sense, alkaline niobates have been considered good candidates as an alternative for this substitution. However, the processing of ceramics with alkaline niobate compositions by classical methods has traditionally shown difficulties in obtaining densities near the theoretical. The classical method for obtaining alkali metal niobate powders by a solid-state reaction between Nb$_2$O$_5$ and alkali metal carbonates involves high temperatures and long reaction times that produces the volatilisation of the alkaline metal, leads to poor compositional homogeneity and provides precursors with large particles. Traditionally, hot-pressing at high temperatures and more recently other pressure- or electric field-assisted methods as the spark plasma sintering of precursors obtained by solid state reaction or other alternative methods are needed to increase the density of the ceramics.

Sodium niobate (NaNbO$_3$), within the series of alkaline-niobate compounds, is antiferroelectric at room temperature. The application of an electric field, or the substitution of Na by Li$^+$ or K$^+$, induce a ferroelectric phase that provides piezoelectric activity, of interest mainly for high frequency devices. Beside the improvement in piezoelectric activity, solid solutions with Li or K based-niobate are often required to improve the processing of the ceramics with respect to the NaNbO$_3$ pure ones. Thus, an important goal in the study of alkaline-niobate ceramics is to solve the difficulties in the NaNbO$_3$ processing, as a way to enhance its piezoelectricity, as well as to allow a further improvement of properties in the NaNbO$_3$-based solid solution ceramics through the control of its microstructure.
Due to the problems in the processing of these ceramics by traditional methods, alternative routes have to be developed to facilitate the production of desired alkaline niobate ceramics in a controllable way. Powder synthesis using evaporation\textsuperscript{12}, sol-gel\textsuperscript{13}, hydrothermal (both simple\textsuperscript{14} or microwave-assisted\textsuperscript{15}), polymeric precursor processes\textsuperscript{16}, microemulsion mediated hydrolytic decomposition of mixed alkoxide solutions\textsuperscript{17} or coprecipitation\textsuperscript{18} have been described in the literature to facilitate the reactions. Densities even near the theoretical one have been achieved by some of these methods\textsuperscript{12}, although piezoelectric characterization of pure NaNbO\textsubscript{3} is not reported in these works. These chemistry based routes require high purity inorganic or organometallic reactants, that are often sensitive to light or humidity and are more expensive than the widely available oxides and carbonates.

Mechanochemical activation has shown to be a successful way to modify the conditions in which chemical reactions usually take place. It has been applied to produce the synthesis of known functional lead-based (PZT\textsuperscript{19}, PMN-PZT-PT\textsuperscript{20}, PZNPT\textsuperscript{21}) and other lead-free\textsuperscript{22,23} piezoceramics, with Bi\textsubscript{4}Ti\textsubscript{3}O\textsubscript{12} composition. A thorough review of the application of this route to ferroelectric materials in general has been recently published\textsuperscript{24}. During the mechanical treatment the particle size of the crystals is reduced and the homogeneity of the mixture is increased. It improves the reactivity of the precursors\textsuperscript{25} and allows the compact of the green pellet to be optimised. The processing of highly dense ceramics\textsuperscript{26-28} is possible in a single thermal process, in which synthesis, grain growth and sintering take place.

Mechanochemical activation has been proved to be an adequate route to synthesis NaNbO\textsubscript{3} compositions. Recently, it was shown\textsuperscript{29} that mechanosynthesis takes place by milling after 40h of a powder mixture of Na\textsubscript{2}CO\textsubscript{3} and Nb\textsubscript{2}O\textsubscript{5}, although the final mixture contains large amounts of amorphous phases and residual traces of the initial
mixtures. The same authors have also studied the effect of different milling energies on the same mixtures. X-ray diffraction (XRD) shows peaks only corresponding to NaNbO₃ composition after 400h of milling, for vials and balls of yttria-stabilized zirconia (YSZ). Time needed for isolate NaNbO₃ can be reduced to 96h by using tungsten carbide vials with heavier balls rotated at higher speed (300 r.p.m against 200 r.p.m. for YSZ media). However, the influence of the mechanical activation on the sintering, microstructure and piezoelectric properties of the ceramics has not yet been determined. This is an important feature because, as will be shown in this work, it is needed, but not sufficient, to have reactive precursors to obtain good piezoelectric ceramics and ways for the ceramic microstructure control are also required.

It is known that the piezoelectricity in ferroelectric ceramics depends on their microstructure (porosity, grain and domain size). The expected reduction of the processing temperature of the ceramics from precursors obtained by mechanical activation should result in a microstructure with reduced grain size, while the pressure applied and high reactivity and specific surface of the nano-sized particles of the powder should maintain porosity values within the limits required for piezoelectric applications. Although in the last years there has been an increasing interest in the topic, the size effects in the functional properties of these alkaline niobate materials are barely found in the literature. Connected to this topic, some interesting features have been recently reported. It was shown that the reduction of the NaNbO₃ particles to submicron size (200-400 nm) induces a non-centrosymmetric phase that improves the piezoelectric properties, as PFM studies of displacement against voltage shows.

In this work, a quantitative study of the microstructure of highly dense submicron-structured NaNbO₃ ceramics, obtained from four different mechanically activated precursors has been carried out. An iterative method was applied to determine
the complex piezoelectric, elastic and dielectric coefficients of the ceramics from complex impedance measurements at the planar resonance of thin ceramic disks. The relation between processing, microstructure and piezoelectric and elastic properties is discussed.

**EXPERIMENTAL PROCEDURE**

NaNbO₃ ceramics have been prepared from precursors obtained by four different routes. The rationale behind the compositions selected and the details of the precursor processing and characterisation can be found elsewhere. Summarizing, stoichiometric mixtures of analytical-grade Nb₂O₅ and sodium reactants (Na₂CO₃, Na₂O or NaOH) were mixed by hand in an agate mortar. The mixture of Nb₂O₅-Na₂CO₃ (hereinafter CNN) was mechanically activated during 30 days, whereas to obtain similar degree of XRD amorphization, the mixtures Nb₂O₅-Na₂O (hereinafter O-NN) and Nb₂O₅-NaOH (hereinafter NOH-NN) were activated for only 7 days. For the fourth precursor, a mixed technique of wet-chemistry plus mechanochemical activation was used. NaOH was dissolved in deionised water and the stoichiometric quantity of Nb₂O₅ was added into the aqueous solution. This suspension was stirred for 4 hours and then dried at 150°C overnight, to obtain the precursor for the mechanical activation process, hereinafter called DOH-NN.

Ceramics were obtained by a single thermal treatment under different conditions. Precursors powders were shaped by uniaxial pressing at 210 kg·cm⁻² as thin disks of 10mm diameter and 2mm thickness, approximately. Disks were isostatically pressed at 2000 kg·cm⁻² and then pressure-less sintering was carried out on a Pt foil at 1200°C for 2h. Alternatively, disks were uniaxially hot-pressed at ~200 kg·cm⁻² in alumina dies and surrounded by alumina powder at temperatures of 900, 1000°C and 1100°C. Hot
pressing time was 2h in all cases, except for the ceramic processed at 900ºC from C-NN precursors (1h).

XRD patterns at room temperature were measured in a D500 Siemens Diffractometer, between 2θ=20º-50º, with an step of 0.05º and counting time of 0.5 seconds per step.

Quantitative microscopy characterisation was carried out with a computerised image analysis and measurement system (IMCO10-KAT386 system, Kontron Elektronic GMBH, 1990) by a procedure explained in detail elsewhere37. Ceramic surfaces were polished and analysed by optical microscopy (Leitz Laborlux 12 ME S/ST), in order to examine the ceramic porosity. The percentage of porosity was characterised by the fraction of the examined area which is occupied by pores. The pore size (P) was characterized by the pore area, since the shape greatly varies from sample to sample. The experimental error of these measurements is ± 0.1 µm². The average and standard deviation of the distributions of pore size were obtained from the linear fitting of the experimental data represented in probability plots38. In these plots, the Ln of the variable being studied is represented against the corresponding cumulative frequency in a probability scale. Linear fitting of the obtained plot provides the size distribution characteristics and demonstrates the single or bimodal character of the distribution37.

Polished ceramic surfaces were etched from 800 or 900ºC to room temperature for the study of the grain distribution. Due to the submicron range of grain size of all ceramics, scanning electron microscopy (960 Zeiss SEM apparatus) was necessary to obtain surface images of the etched ceramics. The ceramic grain size (G) was characterized by the equivalent diameter to a circular shape, a valid approximation for all samples, and calculated from the measured grain area as: D_{eq} = (4·Area/π)^{1/2}. The experimental error of these measurements is ±20 nm. The average and standard
deviation of the distributions of equivalent diameter were also obtained from the corresponding probability plots.

Electroded ceramic samples of diameter \(D\) ~10 mm were grounded to a 0.7-1.0 mm thickness \((t)\) to obtain a sample with a \(D/t\) ratio ~10. Pt paste was sintered at 700°C in the major sample surface electrodes, and then ceramics were poled to saturation in a silicon oil bath at 120-180°C, with fields up to 60 kV/cm. The piezoelectric \(d_{33}\) coefficient was measured in a Berlincourt-meter by the direct piezoelectric effect at 100Hz. Dielectric \((\varepsilon'_{33})\), elastic \((s_{11}^E)\) and piezoelectric complex coefficients \((d_{31})\), as well as electromechanical coupling factors \((K_p)\) corresponding to the radial vibration mode of the disks were measured by an automatic iterative method described elsewhere\(^{39}\), from complex impedance measurements at resonance using a HP4192A impedance analyser.

**RESULTS**

Figure 1 shows the x-ray diffraction patterns of ceramics pressure-less sintered at 1200°C from C-NN, O-NN and DOH-NN precursors. The intensity axis is represented after normalization to 1, so comparison can be directly done. In all cases, a single NaNbO\(_3\) (JCPDS-ICDD file no. 33-1270) phase is obtained, indicating that it is possible to have ceramics in a single sintering thermal treatment, and that no decomposition occurs at that temperature.

Figures 2(a) and 2(b) show the polished surfaces of ceramics sintered at 1200°C from O-NN and DOH-NN precursors, respectively. In the case of C-NN precursors, it is also possible to get single phase ceramics, as Figure 1 shows. However, the density obtained by measurements of dimensions and mass is as low as 45% of the theoretical one, even lower than the green pellet density (63%). It makes impossible to polish the
surfaces, and quantitative microstructure analysis has not been carried out in those types of ceramics.

While polishing, it was observed that the mechanical stability of the DOH-NN and the O-NN ceramics was very poor. Pull-out and cracking occurred even at the smoothest steps of polish (0.1 µm Al₂O₃ emulsion). It makes impossible the use of these sintered ceramics in any further piezoelectric applications. To improve the mechanical stability, and with the aim to preserve a single thermal step process, hot-pressing was applied to obtain mechanically stable and dense ceramics.

Figure 3 shows the x-ray diffraction patterns of ceramics hot-pressed at 900°C(1h) from C-NN, 1000°C from C-NN, O-NN, NOH-NN and DOH-NN, and 1100°C for O-NN and DOH-NN precursors. As for Figure 1, the intensity is represented after normalization to 1. All of them are single phase ceramics, except the C-NN and ONN hot pressed at 900 and 1000°C, respectively. A peak corresponding to a trace of a secondary phase is marked with a circle.

Figure 4 shows the polished surfaces of the hot pressed ceramics. All of them present close porosity, except the hot-pressed at 1000°C from O-NN. This ceramic presents two types of pores, one coming from room among the agglomerates (open porosity) and the other from inside them. The increase in the processing temperature in C-NN to 1000°C and O-NN to 1100°C decrease the porosity. There is an opposite trend for the DOH-NN, in which an increase of hot-pressing temperature from 1000 to 1100°C increases both the porosity and the number of pores, as corresponds to a beginning of a degraded sintering stage

Figure 5(a) shows the probability plots of the pore area of the ceramics shown in Figure 1. The slope of the line for the DOH-NN ceramic is shifted to lower values, meaning wider pore area distribution. From the micrographs and the probability plots,
values of content of porosity, mean value and standard deviation of the pore size
distributions are calculated. The results are shown in Table I. Porosity is slightly lower
for the DOH-NN ceramic (3.0%) than for the O-NN one (4.5%), but it has larger mean
pore size values.

As for the pressure-less sintered ceramics, the Figure 5(b) shows the probability
plot of the pore area of hot-pressed ceramics. The slope of the ceramics hot-pressed at
900°C from C-NN, and 1000°C from O-NN are lower, and the cut with the x-axis
decreases. The slopes are very similar for the rest of the ceramics. Table I also shows
the values of content of porosity, and the mean value and standard deviation of the pore
size distributions obtained from the probability plots of Figure 5(b). The porosity is
higher for the C-NN and O-NN precursors, and it is lower for the DOH-NN one. In this
case, values of porosity as low as 0.40% and 2.10% are obtained by hot-pressing at
1000°C and 1100°C, respectively.

The measurement of the porosity of ceramics from polished surfaces is a
technique successfully used in previous works. For both the pressure-less sintered
and the hot pressed ceramics studied in this work, the linear fit of the probabilistic plot
of the pore area proves that the objects measured are really pores, and not artificial
elements. The content of porosity shown in Table I, measured from the micrographs,
can thus be taken as good, within experimental error.

The problems during polishing for pressure-less sintered ceramics do not appear
in hot-pressed ceramics. The polished ceramics were then etched in order to study the
grain size. The ceramic hot-pressed at 900°C from C-NN precursors was not measured,
because its high porosity (33.7%) reduces its practical interest, in comparison with the
other ceramics studied. Figure 6 shows the aspect of the etched surfaces of the rest of
hot-pressed ceramics. The ceramics processed at 1000°C from O-NN and from DOH10
NN, are inhomogeneous, but the nature of this inhomogeneity is slightly different. In the O-NN case, a clear bimodal distribution with two families of grains that seems to be entirely formed is observed, showing remarkable difference in grain size of the two families. The surface percentage of occupancy of the higher grain size family is 35%, approximately. Ultra-fine intergranular porosity accompanies the family of smaller grain sizes. This ultra-fine porosity, which was not observed at the magnification of Figure 4d, was not taken into account for the measurement of pore size distribution. Such ultra-fine porosity, which is also negligible when calculating the total percentage of porosity is, however, detrimental for the grain connectivity needed for the establishing of a homogeneous electric field in the ceramic.

The DOH-NN ceramic hot-pressed at 1000ºC presents a more complex microstructure comprising three families of objects with characteristic sizes. One of them consists of well formed grains, similar to the grains of the more homogeneous microstructures in Figure 6. Those grains are marked with an asterisk in Figure 6 for DOH-NN 1000ºC ceramic. They occupy approximately 50% of the area of the ceramic surface analysed. Another family of objects consists of clusters, with an internal structure, from which the well formed grains most probably grow as the sintering progresses. Finally, the third family of objects will be the very fine grains in coalescence that constitute the internal structure of the clusters. These are well connected and do not show intergranular porosity. This ceramic presents also an overall fully dense aspect (Figure 4f).

The increase of the hot-pressing temperature to 1100ºC increases the homogeneity in grain distribution for both O-NN and DOH-NN ceramics (Figure 6).

Figure 7 and 8 shows the probability plots of the grain size distributions of hotpressed ceramics. Figure 7 also shows the original distributions of grain size for the
two ceramics with complex microstructures discussed above: the ceramics obtained at 1000ºC from O-NN and from DOH-NN. The good linear fit of the cumulative frequency vs. natural logarithm of the grain size proves the accuracy of the measurement. The measured objects are grains representative of the bulk, and not other artificial objects, appearing for example during thermal etching.

A strong change in the slope of the probability plot is observed for the O-NN 1000ºC ceramic, showing the bimodal character of the distribution of the logarithm of grain size. The linear fitting showed provides the average and standard deviation of the population of smaller grain size, which are shown in Table II. There is not a statistical sample of the population of larger grains at the magnification of Figure 6, and the quantitative analysis was not carried out for this population.

For the also complex microstructure of DOH-NN 1000ºC ceramic, only the distribution that correspond to the population of grains marked with asterisk in Figure 6 is showed and quantitatively characterized in Figure 7 and Table II. The method here used lacks experimental resolution for the measurement and analysis of the population of smaller grains in coalescence. However, their average grain size must be much lower than the one for the population of small grain sizes for the O-NN 1000ºC ceramic, as can be clearly observed by comparison of the corresponding micrographs in Figure 6, i.e., must be well below 100 nm.

The analysis by probability plots of all the other grain size single distributions of hot-pressed ceramics are shown in Figure 8 and Table II. The linear relation between the cumulative frequency and the logarithm indicates again that all of them are lognormal distributions. The slopes move to higher values as the hot-pressing temperature increases, 1000ºC in Figure 7 and 1100ºC in Figure 8, due to the observation of narrower distributions, corresponding to more homogeneous
microstructures. For the sake of comparison with the distributions of the complex microstructures in Figure 7, Figure 8 shows the grain size distribution of the most homogeneous ceramic microstructure obtained, for DOH-NN 1100°C.

Table II also shows the piezoelectric $d_{33}$ coefficients, which depend on the microstructure of the ceramics. The ceramics hot-pressed at 1000°C from O-NN and NOH-NN, which show a combination of a very low average size for the population of small grains (110 nm) and insufficient grain connectivity for the former, and the lowest average value for a single grain size distribution (170 nm) for the latter, are difficult to pole. Their $d_{33}$ coefficients are the lowest (6 pC.N⁻¹) of all the measured ceramics.

Table III shows the complex dielectric, elastic, piezoelectric and electromechanical coupling factors corresponding to the radial resonance mode of poled ceramic disks at room temperature. Only the ceramics with the highest values of $d_{33}$ (Table II) were chosen to be studied. It is noticeable that properties of all ceramics are similar and only small differences among them are observed. The ceramic from C-NN (hot pressed at 1000°C) has the highest piezoelectric coefficient ($d_{31}=-8.7$ pC.N⁻¹) and electromechanical activity ($k_p=14\%$). The ceramic hot-pressed at 1100°C from DOHNN precursor presents the highest elastic properties, with the highest values of planar frequency number ($N_p = 3778$ KHz·mm) and the elastic stiffness ($c_{11}^p = 15.7$ N·m⁻²).

**DISCUSSION**

Figure 1 shows that pressure-less sintering of mechanochemical activated precursors produces single-phase ceramics in a single thermal process, in which synthesis, sintering and grain growth take place. It is due to the reduction by mechanical activation of the thermal barrier necessary to activate the process. However, the density obtained at the same sintering temperature is different for each precursor. For the C-NN
sintered ceramic is lower than the green pellet one because of the well-known effect of the use of carbonates that produce CO₂, which is very volatile, during the thermal treatment\textsuperscript{43}. It hinders the sintering and the closure of porosity. Moreover, the C-NN precursor is less reactive \textsuperscript{36,43} than the other ones, as a higher temperature and a larger time is needed to isolate the phase. The changes in the state of Nb₂O₅ (particle size and microstrains) during milling are negatively affected by the simultaneously activation with NaCO₃ \textsuperscript{30}, reducing the total reactivity of the mixture.

The ceramics obtained by the other precursors show lower porosity (4.5% for ONN and 3.0% for DOH-NN) whilst using lower processing temperatures, than those found in the literature for NaNbO₃ ceramics sintered from solid state reaction precursors (4%, at 1240°C, in two steps\textsuperscript{44}, 5% at 1550°C\textsuperscript{45}) or chemical routes (5%, at 1250°C)\textsuperscript{12}.

As it was said, the problem with pressure-less sintering is that the ceramics reach a high density but they are easily cracked. Although the surface reactivity of the mechanically activated powder particles improves the surface mass transport needed for the sintering process and closure of the interparticle porosity, the resulting bonding between particles seems to be weak, most probably by accumulation in grain boundaries of crystal defects migrated to the surface. The presence of agglomerates may also yield to non homogenous microstructures (in pore and grain size distribution) that could contribute to the cracking. This is critical for piezoelectric applications, in which a poor mechanical resistance affects the long life of the material under vibration or repeated actuation. The use of the hot-pressing solves that problem.

Figure 3 shows that a single phase ceramic can be also obtained in a one-step thermal treatment by hot pressing. Exceptions are C-NN at 900°C or O-NN at 1000°C ceramics. It seems that it is due to the low hot-pressing temperature, not enough for this
not so high reactive precursor to synthesize a single phase. The secondary phase can be indexed as Na$_2$Nb$_4$O$_{11}$ (JCPDF ICPD file no. 72-1694), also observed during the thermal treatment of powders after the mechanical activation of the O-NN precursors$^{36}$. The phase Na$_2$Nb$_4$O$_{11}$ also appears as an intermediate phase when traditional solid state reaction is used. It indicates that O-NN precursors have a lower reactivity than the others, which also affects, as Figure 4 shows, to the porosity content of the ceramics. The main contribution to the porosity comes from the inter-agglomerates space. It increases the porosity (6%), as Table I shows. An increase of the hot-pressing temperature from 1000 to 1100°C makes the secondary phase to disappear (Figure 3) and a decrease in the porosity (from 6 to 3.8%). Consequently, there is also an important reduction in the pore area (from 4.2 to 0.9 $\mu$m$^2$).

DOH-NN precursors give the lowest values of porosity of the hot-pressed ceramics (only 0.40% for 1000°C). A further increase of the hot-pressing temperature to 1100°C produces a slight increase in the porosity (to 2.10%, Table II), due to the decrease in the ratio between sintering and grain growth rate, which is characteristic of a degraded sintering stage.

The residual porosity is reduced with respect to the reported for other alkalineniobate ceramics prepared by hot-pressing of Li$_{1-x}$Na$_x$NbO$_3$ $^6$ (>10% at 1100°C), in which the presence of light Li atoms promotes the mass transport. The residual porosity is also lower than that achieved using other pressure-assisted processing methods, as spark plasma sintering by Wada$^8$ et al. (6% at 1200°C for NaNbO$_3$ ceramics) or Zhang et al.$^9$ (8% at 920°C for Na$_x$K$_{1-x}$NbO$_3$ with x=0.20, where the presence of K also improves the sinterability). In both cases precursors obtained by traditional solid state reaction were used. An important advantage of the use in this
work of lower processing temperatures is that helps preventing the evaporation of the
large amounts of alkaline elements.

The grain size distributions are very inhomogeneous for the ceramics hot-
pressed at 1000ºC from NOH, and much more from O-NN and DOH-NN precursors, as
Figure 6 shows. This is usually related with incomplete sintering stages of various
types. For O-NN precursor it could be due to inhomogeneity by the presence of the
secondary phase observed in Figure 4, which gives place to larger grains observed in
Figure 7. An increase of the temperature solves the problem, and the ceramic hot
pressed at 1100ºC from O-NN precursor has a more homogeneous microstructure as the
comparison of the slopes of the probability plots of the grain size distributions shows
(Figures 7 and 8). The mean grain size and standard deviation of the corresponding
distribution for 1100ºC O-NN ceramic is still relatively small (240 nm and 120nm),
even when compared with the grain size of the ceramics from C-NN (220 nm and 120)
hot pressed at 1000ºC. These results confirm the lower reactivity of the O-NN precursor
with respect to the other ones studied.

The incomplete sintering at 1000ºC from DOH-NN ceramic seems to have
different causes than for O-NN ones, as no secondary phases were found for this
ceramic. It has been reported\textsuperscript{46} that growth in NaNbO\textsubscript{3} during calcination from
homogeneous amorphous polymeric precursors occurs in two steps. Up to temperatures
\(\sim 300ºC\) higher than the synthesis one, a process of neck formation and growth
controlled by surface diffusion occurs, filling the necks among particles. In a second
stage, the densification of agglomerates takes place, and the particle growth increases
substantially in comparison to the crystallite growth. A further increase of temperature
increases the particle size by a process of grain boundary migration. Similarly, during
hot-pressing, the low temperature processes and the densification of agglomerates are
favoured, improving the density of the ceramics. The presence of the clusters in the DOH-NN ceramic hot-pressed at 1000ºC, from which larger grains are grown, seems to indicate that grain boundary migration is not complete. It favours the low porosity achieved with respect to the other precursors, but at temperatures as low as 1000ºC the grain distribution is still bimodal. A further increase of temperature to 1100ºC increases the grain boundary migration and the homogeneity of the grain size.

Table II also shows the $d_{33}$ piezoelectric coefficient of the hot-pressed ceramics. The lower values ($d_{33}=6$ pC/N) obtained for the ceramics processed at 1000ºC from ONN and NOH-NN precursors are a consequence of the low polarizability of those ceramics, already discussed. For ceramics from C-NN and DOH-NN precursors it was possible to pole the ceramics up to saturation at 180ºC with fields of 60 KV/cm, which explains the slightly higher piezoelectric coefficients obtained.

The hot-pressed from C-NN precursor ceramic presents the highest piezoelectric activity, with values of $k_p=14\%$, $d_{31} = -8.7$ pC.N$^{-1}$ and $d_{33}= 40$ pC.N$^{-1}$. These results slightly improve the obtained by Reznitchenko et al.\textsuperscript{44} ($k_p=10\%$) in ceramics processed by classical methods. Values up to $k_p=19\%$ have been obtained by Wada et al.\textsuperscript{8} in spark plasma sintered ceramics (with grain size range of 3-5 $\mu$m). Although all these values are of the same order, the comparison with results in this work is not straightforward, since the method here used is not the IEEE Standard, currently used, but a much more accurate one in the calculation of moderate electromechanical coupling coefficients.

However, it is noticeable that our values are obtained for highly dense submicron-structured ceramics, in agreement with previous works where samples with grain size in the order of 200-300 nm improves the piezoresponse measured by PFM with respect to coarser grain ceramics\textsuperscript{35}. The most remarkable result is the ensemble of piezoelectric and elastic properties ($k_p=12\%$ and $d_{31} = -7.8$ pC.N$^{-1}$ and $d_{33}=38$ pC.N$^{-1}$,
on the one hand, and \( N_p=3252 \text{ KHz.mm} \) and \( s_{11}^E = (9.7-0.03i) \times 10^{-12} \text{ m}^2\cdot\text{N}^{-1} \), on the other hand) that was measured for DOH-NN ceramic hot-pressed at 1000°C with a microstructure characterized by 0.40% residual porosity and a bimodal lognormal distribution of grain size, with average value of 160 nm for the well defined population of higher grain size.

The microstructures obtained in this work also results in good mechanical properties, although comparison with other authors results in this point cannot be made. Slight improvement in the elastic properties by means of the homogenisation of the microstructure in these submicron structured ceramics is also revealed in the study. The ceramic hot pressed at 1100°C from DOH-NN precursor has the highest frequency planar number (\( N_p= 3778 \text{ KHz-mm} \)) and the lower complex elastic compliance (\( s_{11}^E=(6.7-0.008i)\times 10^{-12} \text{ m}^2\cdot\text{N}^{-1} \)) of all the ceramics. The higher homogeneity of the corresponding microstructure is evidenced by the comparison of the grain size distribution of this ceramic in Figure 8 with those of the complex microstructures in Figure 7.

CONCLUSIONS

Ceramics of composition NaNbO₃ have been prepared from precursors obtained by four different routes of mechanical activation. Single phase ceramics are obtained in one-step thermal treatment.

The porosity of the pressure-less sintered is lower than the obtained from solid-state reaction while allows using lower temperatures. However, they have poor mechanical stability that prevents their practical use.

Hot pressing allows obtaining highly dense (with contents of porosity down to 0.4%) and mechanically stable submicron structured ceramics. Homogenous
microstructures with single lognormal distributions and submicron average grain size can be obtained, together with residual porosities of 1.5 and 2.10%, respectively. This submicron structured ceramics is as a consequence of the low particle size of the powder precursors and the low processing temperatures needed when mechanically activated precursors are used. When hot-pressing conditions are not optimized, intermediate stages of sintering are observed and bimodal lognormal distributions of grain size are measured.

A noticeable ensemble of piezoelectric and elastic properties was measured for DOH-NN ceramic hot-pressed at 1000ºC. The observed changes of the functional properties of the ceramics were not high and were related to microstructural characteristics. The highest piezoelectric activity was characterized by values of $k_p=14\%$ and $d_{31} = -8.7\text{ pC.N}^{-1}$ and $d_{33} = 40\text{ pC.N}^{-1}$. The best elastic properties of the ceramics were characterized by values of $N_p= 3778\text{ KHz.mm}$ and $s_{11E}^{E}=(6.7-0.008i)10^{-12}\text{ m}^2\text{N}^{-1}$) and related with the homogeneity of the submicron structured microstructure that shows a single lognormal distribution.

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Figure captions

Figure 1. XRD patterns of pressure-less sintered NaNbO$_3$ ceramics from mechanically activated precursors. [Peak marked with (*) corresponds to Pt, remaining from electrode].

Figure 2. Optical micrographs of the polished surfaces of NaNbO$_3$ ceramics pressureless sintered at 1200ºC from (a) O-NN and (b) DOH-NN precursors.

Figure 3. XRD patterns of hot-pressed NaNbO$_3$ ceramics from mechanically activated precursors. [Peak marked with a circle corresponds to a Na$_2$Nb$_4$O$_{11}$ phase].

Figure 4. Optical micrographs of the polished surfaces of hot pressed NaNbO$_3$ ceramics from mechanically activated precursors: (a) C-NN, 900ºC (1h); (b) C-NN 1000ºC; (c) NOH-NN 1000ºC; (d) O-NN 1000ºC; (e) O-NN 1100ºC; (f) DOH-NN 1000ºC; (g) DOH-NN 1100ºC.

Figure 5. Probably plots of pore size for surfaces of pressure-less sintered and hot pressed NaNbO$_3$ ceramics.

Figure 6. SEM micrographs of polished and thermally etched surfaces of hot pressed NaNbO$_3$ ceramics.

Figure 7. Asymmetric and bimodal grain size distributions and corresponding probability plots, from the corresponding distributions of logarithm of the grain size, for the hot pressed ceramics that present very fine structured areas: (a) O-NN 1000ºC and (b) DOH-NN 1000ºC. Lines of the probability plots (c) correspond to the fitting of the measurable grain size distributions, whose parameters are presented in Table II.

Figure 8. (a) Grain size distributions and (b) corresponding probability plots, from the corresponding distributions of logarithm of the grain size for hot pressed NaNbO$_3$ ceramics: C-NN 1000ºC; NOH-NN 1000ºC; O-NN 1100ºC; DOH-NN 1100ºC. The
grain size distribution of the last ceramic, a single lognormal distribution, is presented for comparison with distributions shown in Figure 7.

TABLE CAPTIONS

Table I. Pore size distribution parameters of pressure-less sintered and hot-pressed NaNbO₃ ceramics. The error quoted is the highest among the experimental and calculation values.

Table II. Grain size distribution parameters of hot-pressed NaNbO₃ ceramics. Values shown for samples marked with asterisk correspond to the measurable grain size distribution of the bimodal type distribution observed in these samples.

Table III. Dielectric, elastic, piezoelectric and electromechanical coupling factors corresponding to the radial resonance mode for hot-pressed NaNbO₃ ceramics.
Figure 3

Intensity (a.u)

2θ(°)

DOH-NN 1100°C-2h
O-NN 1100°C-2h
DOH-NN 1000°C-2h
O-NN 1000°C-2h
NOH-NN 1000°C-2h
C-NN 1000°C-2h
C-NN 900°C-1h

Figure 3
Figure 4
Figure 4
Figure 5

(a) Sintered NaNbO₃ ceramics
- O-NN
- DOH-NN

(b) Hot-pressed NaNbO₃ ceramics
- C-NN 900°C (1h)
- C-NN 1000°C
- NOH-NN 1000°C
- O-NN 1000°C
- O-NN 1100°C
- DOH-NN 1000°C
- DOH-NN 1100°C
Figure 6

- C-NN 1000°C
- NOH-NN 1000°C
- O-NN 1000°C
- O-NN 1100°C
- DOH-NN 1000°C
- DOH-NN 1100°C
Figure 8

(a) Relative frequency distribution of grain size (nm) for C-NN 1000°C, NOH-NN 1000°C, O-NN 1100°C, and DOH-NN 1100°C.

(b) Cumulative frequency plots of Ln (Grain size (nm)) for the same conditions.
# Table I

## PORE DISTRIBUTION

<table>
<thead>
<tr>
<th>Precursor</th>
<th>Sintering Temperature</th>
<th>Mean Pore Size &lt;P&gt; (µm²)</th>
<th>Standard Deviation σₚ (µm²)</th>
<th>Porosity (%)</th>
<th>Number of measured pores Nₚ</th>
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<td><strong>Pressure-less Sintered Ceramics</strong></td>
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<tr>
<td>O-NN</td>
<td>1200°C</td>
<td>0.7±0.1</td>
<td>0.5±0.1</td>
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<td>DOH-NN</td>
<td>1200°C</td>
<td>1.3±0.1</td>
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<td><strong>Hot Pressed Ceramics</strong></td>
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<tr>
<td>C-NN</td>
<td>900°C (1h)</td>
<td>10±2</td>
<td>18±2</td>
<td>33.7±0.5</td>
<td>800</td>
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<td>1000°C</td>
<td>0.7±0.1</td>
<td>0.7±0.1</td>
<td>6.4±0.8</td>
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<td>NOH-NN</td>
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<td>0.4±0.1</td>
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<tr>
<td>O-NN</td>
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<td>0.2±0.1</td>
<td>2.10±0.06</td>
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## Table II

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<th>Precursor</th>
<th>Hot Pressing Temperature</th>
<th>Mean Grain size $&lt;G&gt;$ (nm)</th>
<th>Standard Deviation $\sigma_G$ (nm)</th>
<th>Number of measured grains $N_G$</th>
<th>Piezoelectric coefficient $d_{33}$ (pC/N)</th>
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<tr>
<td>C-NN</td>
<td>1000°C</td>
<td>220</td>
<td>120</td>
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<td>NaNbO₃ ceramic type</td>
<td>C-NN 1000°C</td>
<td>O-NN 1100°C</td>
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<td>Planar coupling factor $K_p$ (%)</td>
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<td>Elastic stiffness $c_{11}^P$ $(10^{-10} \text{ N} \cdot \text{m}^{-2})$</td>
<td>12.8+0.06i</td>
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<td>10.9 + 0.02i</td>
<td>15.7+0.02i</td>
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<tr>
<td>Elastic compliance $s_{11}^E$ $(10^{-12} \text{ m}^2 \cdot \text{N}^{-1})$</td>
<td>8.5-0.04i</td>
<td>7.8-0.01i</td>
<td>9.7 - 0.03i</td>
<td>6.7-0.008i</td>
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<tr>
<td>Piezoelectric coefficient $d_{31}$ $(10^{-12} \text{ C} \cdot \text{N}^{-1})$</td>
<td>-8.7+0.09i</td>
<td>-5.2+0.12i</td>
<td>-7.8 + 0.06i</td>
<td>-7.2+0.03i</td>
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<td>Relative permittivity $\varepsilon_{33}^T$</td>
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