The structure of niobium and tantalum oxides processed by concentrated light flux

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Abstract. The article reports on significant changes occurring in the structure under treatment of Nb$_2$O$_5$ and Ta$_2$O$_5$ ceramics by a concentrated light flux (CLF). The changes comprise, in particular, appearance of micro- and nanostructures, as well as changing type of chemical bonding and coordination number of polyhedrons. These structural changes manifest themselves in the corresponding changes in the physical properties. The influences of the CLF on the Nb$_2$O$_5$ and Ta$_2$O$_5$ ceramics turn out to be different. The number of nanosize structures appearing in the Ta$_2$O$_5$ ceramics is higher, while the changes in the chemical bonding and the coordination of polyhedrons are different. The differences observed by us are explained by differing melting temperatures of Nb$_2$O$_5$ and Ta$_2$O$_5$ and degree of covalence of the chemical bonds Nb–O and Ta–O.

Keywords: niobium and tantalum oxides, structure, concentrated light flux, Raman scattering

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UDC: 535.4, 538.9

1. Introduction

Applications of Nb$_2$O$_5$ and Ta$_2$O$_5$ oxides obtained using conventional ceramic technologies are limited because of a brittle and fragile macro-crystalline structure of these materials. Therefore one can expect that finer micro- and nanoscale structures could improve mechanical properties of the ceramics. In this respect, investigations of the micro- and nanostructures and searching for new methods of materials treatment seem to be urgent [1]. An atomic force microscopy and Raman scattering studies of the influence of concentrated light flux (CFL) on the structure of niobium and tantalum oxide ceramics have been reported earlier. In particular, the Raman spectra have been revealed to be sensitive to interactions of structural units and disordering, thus revealing fine details of irradiation-imposed structural changes, which could not be detected with a common X-ray diffraction analysis [2]. The present study is devoted to further investigations of the effect of CFL on the structure and physical properties of the above ceramics.

2. Experimental methods

Ceramic samples prepared from Nb$_2$O$_5$ and Ta$_2$O$_5$ that contained admixtures with the concentrations less than $5 \times 10^4$ mass per cent were processed using a photo-thermal treatment in the focal

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* The materials of this work have been reported at the 2nd Ukrainian–Polish–Lithuanian Meeting on Ferroelectrics Physics (9–13 September 2012, Lviv, Ukraine).
zone of an optical furnace [3]. A high-performance image analyser Thixomet® and an atomic-force microscope Nano-R2 were used to explore the micro- and macro-structures of the ceramics.

The Raman spectra of the ceramic niobium and tantalum oxides were studied in so-called ‘reflection’ geometry. They were excited using radiation of a MML-100M Y:Al laser (the light wavelength being \( \lambda_0 = 530.5 \) nm) and a Spectra Physics laser (the light wavelength 514.5 nm and the power \( P = 800 \) mW). The spectra were recorded at the room temperature with a modified DFS-24 spectrometer. The parameters of the Raman bands such as the frequency, the intensity and the bandwidth, were retrieved employing a BOMEM GRAMS program.

3. Results and discussion

Processing of the samples by the CLF yields in the conditions, which are far from thermodynamic equilibrium ones. It is performed under fast heating and high thermal gradient in the surface layer (up to hundreds of degrees per millimetre), with the gradients in the layers located deeper in the bulk being two to five times smaller. As a consequence, all of our results should be analysed with considering a possible self-organisation under dissipation of a strong energy flux in the system under examination. A number of new stable and fractal structures considered as attractors could appear. Under the action of energy flux, the structure tends to one of the attractors, its shape and type depending on the fluctuations of different parameters, which are hardly predictable.

![Fig. 1. Micrometer-scale fractal structures appearing in the ceramic Ta₂O₅. Picture plane is parallel to the CLF direction. The pictures are obtained with the Tixomet® image analysis system.](image-url)
The ceramic niobium and tantalum oxides subjected to the CLF treatment contain complex one- or two-dimensional multi-layers that are shaped as 30 nm-thick fibres and 0.5 μm-thick plates and have the linear dimensions about 50 μm. The real structures of several ordering levels observed by us are obviously of a fractal type (see Fig. 1). At the first stage of the process, a competition takes place between the growth centres on the surface and those in the bulk of the material. The structures exhibiting fractal properties are hierarchic and similar at different scales. In other words, if there is no reference scale in the further pictures showing the structures, then it is impossible to distinguish the exact scale.

The nanosize fractal structures of the Ta₂O₅ ceramics processed by the CLF are shown in Fig. 2 and the size distributions typical for these structures are depicted in Fig. 3. The fractal nanosize structures and the corresponding size distributions for the Nb₂O₅ ceramics obtained after the CLF treatment are given by Fig. 4 and Fig. 5, respectively.

The CLF-processed Ta₂O₅ ceramics has a wider range of nanosize structures, when compared with the Nb₂O₅ ceramics (cf. Fig. 3 and Fig. 5). The type and the dimensions of the structure have a strong effect on the physical characteristics of the ceramic oxides of niobium and tantalum. Hence, the CLF-treated oxides reveal the micro- and nanostructures (‘micro-cracks’ and ‘nanocracks’), which are known to suppress the effect of thermal expansion [3, 4]. Both the niobium and tantalum oxides have the phases of variable compositions [5, 6, 7]. The structures characterised by different Nb(Ta)/O ratios and thermal histories may belong to modifications of different symmetry space groups.

At least ten stable or unstable crystalline modifications of Nb₂O₅ have been described till now (see [5, 6, 8, 9]). The most abundant among them are two low-temperature modifications (an orthorhombic (γ) and a hexagonal (β) ones) and a monoclinic high-temperature α-modification, the
melting temperature of which is about 1500°C. The crystallographic parameters of the three modifications are presented in Table 1.

Table 1. Crystallographic parameters of the most abundant modifications of Nb₂O₅ [5, 6, 10, 11].

<table>
<thead>
<tr>
<th>Modification</th>
<th>System</th>
<th>Lattice parameters, Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>α–Nb₂O₅</td>
<td>Monoclinic</td>
<td>a: 2.134</td>
</tr>
<tr>
<td></td>
<td></td>
<td>b: 3.816</td>
</tr>
<tr>
<td></td>
<td></td>
<td>c: 1.947</td>
</tr>
<tr>
<td></td>
<td></td>
<td>c/a: 0.919</td>
</tr>
<tr>
<td>δ–Nb₂O₅</td>
<td>Hexagonal</td>
<td>β: 119.694</td>
</tr>
<tr>
<td>γ–Nb₂O₅</td>
<td>Rhombic</td>
<td>γ: 119.694</td>
</tr>
</tbody>
</table>

Table 2. Crystallographic parameters of the most abundant modifications of Ta₂O₅ [5, 6, 12–14].

<table>
<thead>
<tr>
<th>Modification</th>
<th>System</th>
<th>Lattice parameters, Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>α–Ta₂O₅</td>
<td>Tetragonal</td>
<td>a: 3.81</td>
</tr>
<tr>
<td></td>
<td></td>
<td>b: –</td>
</tr>
<tr>
<td></td>
<td></td>
<td>c: 3.609</td>
</tr>
<tr>
<td></td>
<td></td>
<td>c/a: 0.947</td>
</tr>
<tr>
<td>β–Ta₂O₅</td>
<td>Rhombic</td>
<td>β: 0.63</td>
</tr>
<tr>
<td>(allomeric to γ–Nb₂O₅)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The structures of different Nb₂O₅ modifications are comprised of two types of ReO₆ octahedral blocks, where Re represents a cation. The cross section of one type contains 15 (i.e., 5×3) NbO₆ octahedrons connected to each other at the edges to form two-dimensional layers. The other type contains 12 (i.e., 4×3) octahedrons connected in the same manner to form a layer containing tetrahedral gaps, a part of which is occupied by niobium cations [9]. The structures described above enable formation of phases of slightly different compositions [6, 9] that contain octahedral and tetrahedral blocks connected at the edges or at the tops.

![Fig. 4. Nano-R² image of nanosize structures of the Nb₂O₅ ceramics processed by the CLF.](image)

The tantalum oxide is also polymorphous. The β-modification of Ta₂O₅ is stable up to 1360°C. This phase looks like colourless rhombic crystals (z = 11) of the space group P2mm, with the melting point 1787°C and the density 8.18 g/cm³ [5, 6, 12]. The α-modification of Ta₂O₅ is stable above 1560°C and looks like grey tetragonal crystals (z = 6) of the space group I₄₁/amd (the melting point being 1887°C and the density 8.37 g/cm³). The lattice parameters of the both crystallographic modifications are presented in Table 2. In particular, the both modifications have the same structural units. They contain TaO₆ octahedrons and distorted pentagonal TaO₅ bi-pyramids, forming a three-dimensional framework of bonds of a different type [12–15].
Obviously, the ceramic samples contain different modifications of Nb$_2$O$_5$ and Ta$_2$O$_5$ that belong to different crystalline systems. The X-ray diffraction analysis cannot identify the atomic coordinates of the structure of real samples. Since the calculations of lattice dynamics are also impossible, the Raman spectra remain the only way to identify the real structures of Nb$_2$O$_5$ and Ta$_2$O$_5$ [7, 8, 9]. Due to complexity of our structures and some amount of endless chains, blocks, chain parts, and octahedral and tetrahedral blocks of different crystallographic systems [6, 9–13], the Raman bands of the niobium and tantalum oxides can be correctly assigned only in the case of single-crystalline samples and a certain scattering geometry. Nevertheless, some effects of the CLF treatment of the niobium and tantalum oxides can be revealed by the analysis of the Raman spectra. As an example, Fig. 6 shows the Raman spectra obtained for the CLF-processed niobium oxide ceramics.

The strongest Raman bands are well known to correspond to symmetric vibrations causing the biggest changes in the polarisability of unit cell [2, 9]. So, the most intense Raman bands for the case of niobium oxide ceramics should be assigned to the symmetric vibrations. The Raman spectra of the Nb$_2$O$_5$ samples contain four groups of intense lines separated by wide energy gaps, thus implying that the corresponding vibrations either do not interact or their interaction is negligible.

The bands around 30–120 cm$^{-1}$ correspond to the vibrations of octahedrons as a whole [14]. The bands in the region of 150–400 cm$^{-1}$ can be referred to the vibrations of cations located inside
the octahedrons and tetrahedrons. The bands near 500–1100 cm\(^{-1}\) specify the vibrations of oxygen framework. Notice that the bands located in the region of 500–800 cm\(^{-1}\) should refer to stretching oxygen framework [14–17]. The bands at 850–910 cm\(^{-1}\) are associated with stretches of the oxygen bridges of octahedrons and tetrahedrons. The spectral band located at 548 cm\(^{-1}\) could be attributed to bending of Nb–O–Nb bonds. Finally, the bands around 500–1000 cm\(^{-1}\) may be ascribed to stretching Nb–O bonds.

![Raman spectra of tantalum oxide ceramics measured before (curve 1) and after (curves 2 to 5) the CLF processing with different relative intensities: \(I_0 = 0\) (2), 0.26 (3), 0.5 (4), and 1.0 (5).](image)

Fig. 6 reveals that the parameters of different Raman bands respond differently to the CLF treatment of the sample. We assume that the intense bands at 55 and 118 cm\(^{-1}\) correspond to symmetric vibrations of the octahedrons and tetrahedrons. At the same time, the bands located in the region of 150–400 cm\(^{-1}\) should be referred to symmetric translational vibrations of niobium ions located in the octahedral and tetrahedral gaps of the structure.

The relative intensities of most of the Raman bands for the CLF-processed Nb\(_2\)O\(_5\) ceramics increase with increasing CLF intensity. For example, one can observe increasing intensities of the bands located at 118 cm\(^{-1}\) (symmetric librations of tetrahedrons) and 1002 cm\(^{-1}\) (Nb–O stretching) under such conditions. Only the intensity of the band located at 55 cm\(^{-1}\) (symmetric librations of octahedrons) remains unchanged. Therefore breaking of Nb–O–Nb bonds inside the octahedrons and tetrahedrons induced by the CLF manifests itself as a growing intensity of the band referred to stretching Nb–O bonds at the free tops. As a result, the structure of the niobium oxide ceramics is broken up into islands of bonded octahedrons and tetrahedrons.

An increase in the number of isolated octahedrons and tetrahedrons occurring with increasing CLF intensity is caused by several processes that accompany each other. First, we have fast (around 10 s) melting of the ceramic niobium oxide under the CLF treatment. Second, a partial dissociation in the liquid phase takes place. And third, fast crystallisation under the conditions far from thermodynamic equilibrium should happen after leaving the CLF zone, the latter apparently leading to formation of an island structure. Together with the appearance of micro- and nanostructures under the above treatment, this type of structure contributes to the changes seen in the mechanical characteristics and it should suppress thermal expansion of the samples [4].

As an example, Fig. 7 demonstrates that the biggest changes are detected for the line vibrations [18]. It is also seen from Fig. 7 that the most substantial changes in the Raman spectra for the ceramic tantalum oxide are observed in the region of 450–1000 cm\(^{-1}\), which corresponds to stretching oxygen framework [18]. A weak band at 903 cm\(^{-1}\) is observed in the Raman scattering originated from that part of the sample surface which has been directly irradiated. It should correspond to stretching Ta–O bonds at the free tops of tetrahedrons. On the one hand, the band located at 980 cm\(^{-1}\) and assumed to refer to Ta–O stretching at the free tops of octahedrons is absent (see...
curve 5 in Fig. 7). This suggests a structure of long octahedron chains and fragmented tetrahedron chains. On the other hand, the Raman scattering from the sample surface irradiated with a lower relative intensity (~ 0.5 – see curve 4 in Fig. 7) reveals a weak band at 981 cm\(^{-1}\) that corresponds to stretching Ta–O bonds in the isolated TaO\(_6\) octahedrons [13, 18]. The band at 981 cm\(^{-1}\) indicates to a presence of shorter isolated fragments of TaO\(_6\) octahedron chains. Moreover, Ta–O–Ta bridges at the ends of these chains are destroyed in those places of the surface where the CLF intensity is substantially lower [18–20].

It is also interesting that the band ascribed to stretching Ta–O bonds at the free tops of octahedrons is absent in the Raman scattering from the sample surface directly irradiated by the CLF (\(I_r = 1\)), whereas practically all the rest of the Raman bands present are the narrowest (Fig. 7). The facts observed by us suggest that the direct CLF irradiation imposes polymerisation of pieces of the octahedron chains, which are linked into extended structures of oxygen polyhedrons [13, 18–20]. Of the reasons for the Raman spectra of the CLF-processed ceramic niobium and tantalum oxides to be different, one can remind a higher melting temperature for TaO\(_2\) and a higher degree of covalence of Ta–O bonds, when compared with that characteristic of Nb–O ones.

4. Conclusions

The most important and general conclusion drawn on the basis of our results can be summarised as significant changes occurring in the structure of ceramic NbO\(_2\) and TaO\(_2\) owing to the CLF. This concerns different ordering levels, including the appearance of micro- and nanostructures, the type of chemical bonding, and the polyhedral coordination. The structural changes in the niobium and tantalum oxides manifest themselves in the physical properties, e.g. mechanical characteristics and thermal expansion [4]. The effects of the CLF treatment on the ceramic NbO\(_2\) and TaO\(_2\) are rather different: the number of the nanosize structures in TaO\(_2\) exceeds that of the NbO\(_2\) ceramics and the changes in chemical bonding and polyhedral coordination are distinct. The differences observed by us can be explained by differing melting temperatures and degrees of covalence of the chemical bonds Ta–O and Nb–O. A higher melting point for TaO\(_2\) decreases the time needed for mutual transformations of the liquid and solid phases after the sample is removed from the zone of irradiation. Quite likely, this should favour formation of the nanosize structures.

References


Анотація. У роботі виявлено суттєві зміни структури внаслідок обробки керамічі Nb$_2$O$_5$ і Ta$_2$O$_5$ концентрованим потоком світла. Ці зміни полягають у нові мікро- і наноструктур, зміні типу хімічного зв’язку, координаційного числа та ін. Зміни структури виявляються у змінах фізичних властивостей. Вплив концентрованого потоку світла на кераміку Nb$_2$O$_5$ і Ta$_2$O$_5$ різний. Зокрема, кількість нанорозмірних структур, що з’являються в кераміці Ta$_2$O$_5$, вища, а зміні хімічного зв’язку і координаційного числа – різня. Ці відмінності пояснюються відмінностями температур плавлення кераміки Nb$_2$O$_5$ та Ta$_2$O$_5$ і ступені ковалентності хімічних зв’язків Nb–O і Ta–O.