Effect of small mobile cations on molybdenum-borate glasses

Abstract

Lithium borate glasses modified with molybdenum oxide according to the formula: xMoO₃-(1-x)(0.25 Li₂O 0.75 B₂O₃) prepared by the melt quench technique were studied. Density, differential thermal analysis, FTIR and impedance spectroscopic were used to analyze these systems. It was confirmed that the glassy matrix of these systems was based on the MoO₃ tetrahedral units and on B₂O₃ and BO₃ units. The results explained how the stability of a borate matrix that hosted molybdenum ions was affected by the presence of small mobile ions (lithium ions in the present case). It was found that concentration of molybdenum oxide was restricted by strong interactions between the charged molybdenum structures and the mobile ions due to the presence of order forces that promote the formation of crystalline structures.

Keywords: molybdenum, oxide glasses, electrical properties, borate glass stability

Introduction

In the field of glass technological applications it is fundamentally important to understand the influence of the structure on the glass properties. It seems curious to speak about the structure of glasses, but it is acknowledged that the three-dimensional order of components is not random in the case of oxide glasses. Modified glasses with transition metal ions are interesting in the spectroscopic field because their properties make them suitable for optic fibers, Fresnel lenses, etc. Binary glass systems based on Na₂B₅O₉ with different transition metal oxides have already been thoroughly studied elsewhere. Nowadays, these glasses are of interest for laser and infrared detection techniques. B₂O₃ is a basic glass former because of its higher bond strength, lower cation size and small heat of fusion. Also, a large set of crystalline borates as: BaB₄O₇, BiB₂O₅, SrB₂O₅, CsLiB₄O₇, LiNaB₄O₇, LiB₂O₅ have been extensively studied because of their particular physical properties in non-linear optical devices. Li₂O-B₂O₃ system also yields Li₂B₂O₅ compound which is technologically important due to its surface acoustic wave (SAW) and its piezoelectric and pyroelectric properties. Although the optical properties of Li₂B₂O₅ have been studied in detail, the literature on the electric properties is limited. Taking into account that molar boron oxide does not crystallize even when it is cooled at a slow rate, it can be considered as the best glass maker because it is a network former in which its coordination can be either threefold and/or fourfold. However, the glassy network features given by this oxide depends on its composition (i.e. on the glass former oxides and/or modifiers oxides incorporated) and the melting conditions applied.

Experimental

Materials were prepared by melting a mixture of weighed amounts of reagents-grade chemicals, namely MoO₃, B₂O₃ and Li₂CO₃ in a platinum crucible at a temperature of 950°C for 2 hours. To ensure homogenization and the complete gas liberation of the carbonate decomposition, the mixture was stirred several times. Molten glass was poured in drops onto a preheated aluminum plate at 200°C. The nominal composition of the prepared glasses is given in Table 1. The electrical conductivity of these materials was determined by a.c. impedance spectroscopy with an Agilent 4284A LCR meter in the frequency range from 20Hz to 1MHz in a temperature domain from 60°C to (Tg – 15) °C. A glassy disk with parallel faces was obtained by polishing a drop. After that, each face of the cylinder was coated with...
a fine layer of silver paint in order to have proper electrical contact. Differential Thermal Analysis (DTA) measurements were carried out applying a heating rate of 10 K min⁻¹, in a temperature range of 298 K to 1273 K on a Rigaku Thermoflex TG 8110 associated with Thermal Analysis Station TAS 100. X-ray powder (XRD) was performed with a PW1710 BASED instrument in continuous scan mode with a copper anode and 45 KV–30 mA for the tension and electrical current generator respectively. Samples were exposed to the Cu Kα radiation (λ = 1.54 Å) at room temperature in the 20 range: 10°–60°. For FTIR spectra, each sample was previously ground in an agate mortar to obtain a very fine powder. A semi-quantitative dispersion of each powdered sample in Nujol was registered on a Nicolet Nexus FTIR instrument, in the 2000–400 cm⁻¹ range, at room temperature, using KBr windows. Density measurements were taken by the Archimedean method using 2-propanol alcohol as secondary displacement medium.

The average of the three independent measures is reported.

### Table 1 Nominal composition of prepared materials.

<table>
<thead>
<tr>
<th>Glass code</th>
<th>Li₂O (mol%)</th>
<th>MoO₃ (mol%)</th>
<th>B₂O₃ (mol%)</th>
</tr>
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<tbody>
<tr>
<td>Mo-0</td>
<td>25</td>
<td>0</td>
<td>75</td>
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<tr>
<td>Mo-1</td>
<td>22.5</td>
<td>10</td>
<td>67.5</td>
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<tr>
<td>Mo-2</td>
<td>20</td>
<td>20</td>
<td>60</td>
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<tr>
<td>Mo-3</td>
<td>17.5</td>
<td>30</td>
<td>52.5</td>
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</table>

### Results

Figure 1 shows the XRD patterns. Samples without or with a low MoO₃ content evidence two different deviations from the baseline, above 20° and 40°. The absence of sharp peaks in Mo-0, Mo-1, and Mo-2 allows confirming that samples are glasses. However, Mo-3 shows several sharp peaks due to the formation of crystalline structures. In this light, it is inferred that the formation of a borate glass is facilitated by the incorporation of MoO₃ in low quantities (i.e. less than 30 mol%) when small mobile ions like Li⁺ are present in a proportion around of 20 mol%. Figure 2 displays glass transition temperature as a function of molybdenum oxide content which passes through a minimum value at 10 mol% of MoO₃. In spite of the fact that Mo-3 is not completely glassy, the T_g of the glass fraction can be identified. Therefore, molar volume (V_m) and oxygen packing density (OPD: the number of mol of oxygen atoms per dm³ of glass) calculated from density values are plotted in Figure 3. As a result, it should be assumed that the increment in the content of MoO₃ involves a compactness of the glassy matrix. According to the literature, borate glasses are built fundamentally on polyhedra of triangular coordination and the addition of alkali oxides as Li₂O forces some of the boron atoms to change from trigonal to tetrahedral coordination and, with concentrations of alkali oxide higher than 20%, NBO (no-bridging oxygen) units are formed. In the system studied in the present work, the lowest glass transition temperature corresponds to the Mo-1 in which the alkali oxide (Li₂O) content is 22.5 mol% and 10 mol% of MoO₃. The initial incorporation of MoO₃ makes the matrix less stable. Nevertheless, higher MoO₃ content (x>0.1) helps to stabilize the matrix and the systems, even in the presence of a moderate alkali oxide concentration, and increases their T_g value. These results explain that the connectivity among boron and molybdenum polyhedra is favorable when the mol MoO₃ / B₂O₃ ratio is not too large where the presence of the alkali oxide causes a minimum perturbation in the 3D connectivity.

From the results in Figure 3 we learn that when a small quantity of molybdenum oxide is incorporated it cuts drastically the glassy matrix (ΔT_g ≈ 75.12 °C, (Table 2). After that, as the MoO₃ concentration increases, the matrix compactness continuously decreases. Molybdenum oxide raises around 20% the glass density value (Table 2) because its molar mass is greater than that of lithium and borate oxides together. However, the molar volume also rises when the molybdenum oxide content augments and as a result, the glassy matrix becomes less compact because of the expansion of its structure. Therefore, the oxygen packing density (OPD) decrease can be interpreted in terms of the formation of a less rigid matrix and resulting in a loosely packed glass compared to the binary one. This reduction agrees with previous results obtained in the case of tellurite glasses. A less compact glassy matrix could facilitate the displacement of the mobile ions (Li⁺). Figure 4 shows the samples infrared spectra. According to the literature, the intensities of the bands from 1000 to 1200 cm⁻¹ (A) decrease as MoO₃ content increases because the substitution of B₂O₃ by MoO₃ diminishes the number of BO₄ groups and induces the formation of tetrahedrically coordinated BO₃ units (B). Another band lies close to 700 cm⁻¹ (C), which is attributed to the coupling of MoO₃ in chain-like arrangements. The studied samples conductivity temperature dependence below the T_g.
shown in Figure 5. These data fit well in the general Arrhenius-type formula:

$$\sigma = \sigma_0 e^{-\frac{E_a}{kT}}$$  \hspace{1cm} (1)

Where $\sigma$ is the pre-exponential term, $E_a$: the activation energy for the process involved, $T$: the absolute temperature and $k$: the Boltzmann constant. Since only one slope is obtained, we assume that a single conduction process takes place. Despite having of molybdenum in sample Mo-1 to Mo-3, we can assume that the conductivity measured is mostly ionic as it is inferred from the Ea values. Nevertheless, the literature leads us to believe that the conductivity due to the presence of molybdenum in the matrix could be the result of a polaron hopping mechanism. However, if such was the case, it would not be the dominant mechanism as we learn from Figures 6A & Figure 6B. These figures show the conductivity values at 523K, the mean Mo-Mo distance and activation energy as a function of molybdenum oxide fraction. Imre et al.23 showed that in single-alkali borate glasses the dc conductivity reveals a power-law behavior with an increasing lithium oxide content for $x > 0.1$.24 It has been shown that in glasses based on vanadium-tellurite oxide, the incorporation of molybdenum causes a diminution of the polaron conductivity.25 Those results are related to relative atomic distances between molybdenum ions. The conductivity decreases upon increasing the oxide molybdenum content might be explained by the presence of MoO$_2$ as network former.26 In spite of the probable presence of molybdenum ions in different oxidation states (ox.-red. couples), the conductivity could still be ionic and even if the polaronic conductivity was present it would be negligible as it is single inferred from the Arrhenius behavior in Figure 5 and the activation energy values obtained from that. Taking into account that the average distance between two molybdenum ions was estimated, using the experimental density values and elementary chemical considerations, as:

$$d = \left(\frac{M_2}{3\rho N_A}\right)^{1/3}$$  \hspace{1cm} (2)

where $M_2$ is the sample molar mass, $\rho$ is the density, $N_A$ is Avogadro’s number and $x$ is the molar fraction of the ion of interest, it is evident from the results plotted in Figure 6B that the individual hops of the electron between active sites for polaronic hopping transport (red-ox sites) are getting closer to one another, as the molybdenum content increases. In this boron oxide glass the dc conductivity reveals a power-law behavior with an increasing lithium oxide content for $x > 0.1$.24 It has been shown that in glasses based on vanadium-tellurite oxide, the incorporation of molybdenum causes a diminution of the polaron conductivity.25 Those results are related to relative atomic distances between molybdenum ions. The conductivity decreases upon increasing the oxide molybdenum content might be explained by the presence of MoO$_2$ as network former.26 In spite of the probable presence of molybdenum ions in different oxidation states (ox.-red. couples), the conductivity could still be ionic and even if the polaronic conductivity was present it would be negligible as it is single inferred from the Arrhenius behavior in Figure 5 and the activation energy values obtained from that. Taking into account that the average distance between two molybdenum ions was estimated, using the experimental density values and elementary chemical considerations, as:

$$d = \left(\frac{M_2}{3\rho N_A}\right)^{1/3}$$  \hspace{1cm} (2)

we analyze the FTIR results presented in Figure 4, the incorporation of MoO$_3$ in the boron oxide glassy matrix provokes the microscopic changes which tends to form a tetrahedrally connected framework and the more connected the matrix, the less available sites for the lithium ion hopping (i.e. a small number of NBO: non-bridging oxygen) it originates. To reinforce our hypothesis on the interactions among the mobile ions (lithium ions) with molybdenum content, we studied the same composition but replacing the MoO$_3$ by WO$_3$ and by CeO$_2$. Figure 7 shows the comparison of the electrical conductivity of these systems. Rao et al.16 studied the ESR spectroscopic properties of Li$_2$O -ZnO-B$_2$O$_3$ glass with different contents of MoO$_3$ (from 1 to 5 mol %) prepared by melt quenching. They found that the molybdenum ions, expected to exist mostly in the Mo$^{6+}$ state when the MoO$_3$ concentration increases beyond 5.0mol. Additionally, IR and optical absorption spectra of paramagnetic ion in such glasses reveal that the site symmetry of the transition metal ion is that of an elongated octahedron. B$_2$O$_3$ glasses are good glass formers and in them boron (B$^{3+}$) ions are present as triangular coordinated polyhedra of oxygen forming the main structural units: BO$_3$ triangles of three member rings connected by B–O–B linkage named boroxol. Those rings are planar with a bond length of 1.36Å.28 BO$_3$ tetrahedral units have a B–O bond length of 1.47Å.29 Borate glasses IR spectra of boroxol rings characteristic absorption is at 806 cm$^{-1}$.29 Previous structural studies of borate glasses have shown that the addition of a network modifier oxide provokes a conversion of the BO$_3$ units to BO$_4$ tetrahedrals incorporated in complex cyclic groups (called diborate, triborate, tetraborate or pentaborate) and a formation of NBO atoms.11,30 Therefore, the absence of the band of boroxol ring (806 cm$^{-1}$) in a borate glass suggests that the glass consists of randomly connected BO$_3$ and BO$_4$ units.32

The conversion of trigonal BO$_3$ structural units into BO$_4$ tetrahedral unit observed in FTIR absorptions results in the increase of NBOs which facilitates the lithium ion hopping. However, because the Mo$^{6+}$ cation has a high field strength, it induces an ordering effect on the oxygen ions. Therefore, if a silicate or borosilicate glass modified with molybdenum oxide is combined with alkali oxide, it tends to form crystalline molybdates. It has been shown that the molybdenum in those glassy systems appears as unconnected tetrahedral MoO$_4$ units within an alkali rich glass causing the glass network depolymerization (i.e., the oxygen atoms of MoO$_4$ units are not bound to silicon or boron atoms). The similarity between the local environment of Mo$^{6+}$ ions in glasses and in molybdate crystalline phases may explain the low solubility of molybdenum and their tendency to crystallize during the slow cooling of melts or heat treatments. The amount of alkali cations able to form NBO diminishes with the increase of molybdenum concentrations showing that molybdenum keeps alkali ions in its surrounding.11 Therefore, as the glass compositions have mostly the same short-range structure as the crystalline one, it is clear that the intermediate-range structure is strongly dependent on the mobile ions present.11 Figure 8 shows that the permittivity real part increases at every temperature when frequency decreases. The real part of the permittivity goes down at every frequency as the content of MoO$_3$ rises. The comparative diminution is particularly significant at low frequencies due to the lower ion polarization at the interface, not only because the nominal concentration is slightly lower but also because a large number of the mobile ions are trapped around molybdenum charged units. Therefore, in the absence of molybdenum, when temperature rises, the dielectric dispersion shifts towards higher frequencies, the environment around Li$^{+}$ is changed and it has an influence on the mobility of Li$^{+}$ ions.
Table 2 Glass transition temperature, density, molar volume and oxygen packing density (OPD).

<table>
<thead>
<tr>
<th>Glass code</th>
<th>Tg [°C]</th>
<th>Density [g.cm$^{-3}$]</th>
<th>Molar volume [cm$^3$.mol$^{-1}$]</th>
<th>OPD [mol.dm$^3$]</th>
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<td>Mo-0</td>
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<td>Mo-3</td>
<td>539</td>
<td>2.77</td>
<td>30.69</td>
<td>86.35</td>
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</table>

Figure 3 Molar volume and oxygen packing density as function of MoO$_3$ content.

Figure 4 FTIR spectra of studied samples.

Figure 5 Arrhenius plot for the electrical conductivity.

Figure 6 Conductivity at 523 K and activation energy (A), mean Mo-Mo distance and activation energy (B), both as function of xMoO$_3$ content.

Conclusion

In the present work, we showed that the stability of a borate glass when acting as a host of molybdenum oxide is influenced by the presence of small mobile ions. We studied the interaction between lithium ions with the molybdenum-borate glass which limits molybdenum concentrations before the formation of crystalline structures. We observed in the borate glass x MoO$_3$, (1-x)[0.25Li$_2$O 0.75 B$_2$O$_3$] prepared by the melt quenching technique that the network structure was based on MoO$_6$ octahedral units and on BO$_3$ and BO$_4$ units. However, the charged molybdenum structures interact strongly with those mobile ions present in the matrix and induce order forces that promote the formation of crystalline structures. Such devitrifications are the reason of the aging of the glasses studied here.

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Conflicts of interest

Authors declare that there is no conflicts of interest.
References