1. Introduction

Phosphate compounds are interesting glass formers and are largely used for their specific properties with respect to other classical glasses such as silicate, borate or germanate glasses. Particularly, they are well known for their small liquidus viscosity, softening temperatures, large UV transparency and high solubility for other glass modifiers or intermediaries such as alkaline, rare earth or transition metal compounds. Another interesting behavior of phosphate glasses is their ability to incorporate fluoride compounds without reduction of the glass forming tendency. These vitreous materials are known as fluorophosphate glasses and are of great interest for rare earth luminescence since the fluoride compounds strongly reduce the phonon energy of the glass host. These glass samples were heat-treated near the crystallization peaks and the X-ray diffraction measurements pointed out that the dominant phase which precipitates from the glass samples containing 40% and 50% of PbF₂ is the lead fluorophosphate phase Pb₅F(PO₄)₃, whereas the sample containing 60% of PbF₂ exhibits a preferential bulk crystallization of cubic lead fluoride β-PbF₂.

Keywords: glasses, phosphate, lead fluoride

2. Experimental Part

Investigated compositions were prepared from the starting compounds NH₄H₂PO₄ 48% in P₂O₅, Na₂CO₃ 98%, WO₃ 99,9%, PbF₂ 99,9%, Er₂O₃ 99,99% and Yb₂O₃ 99,99%, all from Aldrich. The starting powders were weighted using an...
analytical powder balance and ground in an agate mortar. The resulting powder was transferred to a covered platinum crucible, heated at 400 °C for 1 hour to remove residual moisture and absorbed gases, at 600 °C for 1 hour to promote NH₄H₂PO₄ and Na₂CO₃ decomposition and at 850 °C for 10min for melting of these starting powders. The materials were kept at this temperature for only 10min to minimize fluoride losses and the melt was manually homogenized after 5min and 10min. Finally, the melt was poured in a steel mold preheated 20 °C below Tg and annealed at this temperature for 4 hour before cooling to room temperature inside the furnace. DSC curves were obtained on bulk and powder glass samples of 10mg in aluminum pans between 160 °C and 470 °C at 10 °C/min under N₂ atmosphere. These thermal analyses were obtained using a DSC 200 F3 Maia calorimeter from Netzsch. X-ray diffraction measurements were performed on powder samples using a RigakuUltima IV diffractometer working at 40KV and 30mA between 10° and 70° in continuous mode of 0,02°/s. The crystalline phases were identified according to X-ray powder diffraction patterns (PDF file)³⁵.

3. Results and Discussion

The glass forming ability of Er³⁺/Yb³⁺-codoped compositions in the ternary system (100-x)[0,6NaPO₃-0,4WO₃]-xPbF₂ were investigated by melt-quenching and it was found that glasses can be obtained from x=0 to x=50 whereas the sample with x=60 crystallized under these cooling conditions. The molar NaPO₃/WO₃ ratio of 3/2 was chosen because of the known high thermal stability of the binary glass 60NaPO₃-40WO₃ and its suspected high ability to dissolve lead fluoride. In fact, this assumption could be verified since PbF₂ incorporations as high as 50 mole% were achieved for the rare-earth codoped compositions and 60 mole% for the undoped composition as seen in Table 1 and Figure 1. Samples 0PbEY and 10PbEY also exhibit a strong dark color whereas samples containing 20% or more of PbF₂ were transparent. This visible absorption has already been described in the binary system NaPO₃-WO₃ to be due to tungsten reduction from W⁶⁺ to W⁵⁺ in WO₆ clusters³⁶. Visible and near infrared absorption is related with both electronic d-d transitions of W⁵⁺ and polaron transitions between W⁶⁺ and W⁵⁺, resulting in a very broad absorption band between 500nm and 1300nm. This partial tungsten reduction taking place inside WO₆ clusters is known in tungsten oxide crystalline materials and amorphous thin films and is commonly attributed to oxygen loss in order to reach lower free energy. In our case, it is suggested that a similar oxygen loss occurs during melting but is only possible in high WO₆-concentrated glasses where tungsten oxide clusters are formed. Then, it is assumed that increasing PbF₂ contents progressively breaks the covalent network and W-O-W bonds from these WO₆ clusters and hinders the tungsten reduction. Since the main objective of this work is the selection of promising glass compositions for lead fluoride precipitation, the most PbF₂ concentrated glass samples were chosen for further characterizations (40PbEY, 50PbEY and 60Pb). Thermal properties were investigated by DSC to determine the characteristic temperatures and thermal stability in function of the composition as shown in Figure 2 and Table 1. The first general information extracted from these results is that the thermal behavior of the glass samples is strongly dependent of the lead fluoride content. The glass transition temperature Tg decreases from 270 °C to 189 °C, which is in agreement with the expected modifier behavior of lead fluoride in the tungsten phosphate network. In fact, incorporation of fluoride compounds in covalent vitreous networks usually breaks the bridging bonds which are converted to terminal bonds. In our case, it is assumed that terminal P-F and/or W-F bonds are formed and result in a strong decrease of the network connectivity. Thus, the glass transition temperature as well as thermal stabilities decrease since this less connected network is less viscous and tend to crystallize. Another important result extracted from the DSC curves is the identification of different crystallization peaks for each sample. Glass 40PbEY exhibits a very weak exothermic event at 470 °C and an intense peak centered at 530 °C. The baseline shift around 375 ºC could not be related with a crystallization event since heat treatment

| Sample | Molar composition | Tg         | Tx₁ | Tc₁ | Tx₂ | Tc₂ | Tx₁-Tg |
|--------|-------------------|------------|-----|-----|-----|-----|--------|
| 40PbEY | 35,64NaPO₃-23,76WO₃-40PbF₂-0,5Yb₂O₃-0,1Er₂O₃ | 270 °C | 456 °C | 470 °C | 509 °C | 531 °C | 186 °C   |
| 50PbEY | 29,64NaPO₃-19,76WO₃-50PbF₂-0,5Yb₂O₃-0,1Er₂O₃ | 237 °C | 312 °C | 325 °C | 386 °C | 397 °C | 75 °C    |
| 60Pb   | 24NaPO₃-16WO₃-60PbF₂ | 189 °C | 235 °C | 250 °C | 285 °C | 301 °C | 46 °C    |

Figure 1. Glass samples obtained in the ternary system NaPO₃-WO₃-PbF₂.
at 400 °C didn’t induce any crystallization as shown above in this work. Sample 50PbEY presents to main crystallization peaks at 325 °C and 396 °C as well as a clear melting event at 432 °C. Finally, the DSC curve of undoped sample 60Pb is characterized by a first crystallization event at 250 °C, a second exothermic peak at 301 °C with a shoulder at 327 °C and another weak exothermic event at 396 °C. At higher temperatures, the baseline rise is attributed to fluoride evaporation. This distinct crystallization behavior with composition suggests that different crystalline phases can be obtained in each sample under specific annealing conditions. For these reasons, heat treatment were performed at specific temperatures for each sample and the X-ray diffraction measurements of these treated samples are resumed in Figure 3 for composition 40PbEY, Figure 4 for composition 50PbEY and Figure 5 for composition 60Pb. As explained above, the apparent DSC baseline change for sample Pb40EY is not related with any crystallization event since this sample heat-treated for 30 min at 400 °C remains completely amorphous (Figure 3). On the other hand, heat treatment at 520 °C (above the second onset of crystallization $T_x$) for 4 hours promotes crystallization of the lead fluorophosphate phase Pb$_5$F(PO$_4$)$_3$. Other weak diffraction peaks observed in the diffraction pattern could not be identified (labelled * in Figure 3) and may be related with precipitation of another crystalline phase identified by DSC around 470 °C. X-ray diffraction patterns of sample 50PbEY heat-treated at 300 °C (onset of the first crystallization event) for 2 hours and 390 °C (onset of the second crystallization event) for 4 hours are presented in Figure 4. The first heat-treated sample is mainly amorphous since the most intense event is the diffraction halo. A very weak diffraction peak centered at 30,5° is also observed. For the other sample heat-treated at higher temperature, intense diffraction peaks are observed without any residual diffraction halo, indicating the high crystallinity of this sample. Most of these diffraction peaks were attributed to the lead fluorophosphate phase Pb$_5$F(PO$_4$)$_3$, as illustrated in Figure 4. Other weak diffraction peaks (identified by * in Figure 4) were also observed but could not be attributed to any known phase containing the elements Pb and F.
inferred that the first crystallization peak is mainly related to the high temperature crystallization event, suggesting that the main diffraction peak of PbF2(P04)3 is centered at 30.5°, suggesting that the weak diffraction peak observed for the sample annealed at 300 °C is also due to a early stage of PbF2(P04)3 crystallization. Thus, it can be inferred that the first crystallization peak is mainly related with precipitation of PbF2(P04)3. These results point out that glass compositions 40PbEY and 50PbEY are not promising for luminescent glass-ceramics since PbF2(P04)3 is not a suitable crystalline host for rare earth ions due to high phonon energy of phosphate compounds. X-ray diffraction patterns of undoped sample 60Pb heat-treated at 225 °C (onset of the first crystallization event) for 1 hour and 300 °C (onset of the second crystallization event) for 4 hours are presented in Figure 5. For this sample, it could be determined that cubic lead fluoride is precipitated under heat-treatment in both cases with a higher crystallinity of the sample for higher heat-treatment temperatures. In addition, results shown in Figure 5 demonstrated that no other crystalline phase is precipitated under these conditions, suggesting that β-PbF2 growth can be easily controlled in the glass sample in order to achieve the desired crystallite size and distribution. These X-ray diffraction measurements are helpful to attribute the first crystallization peak to cubic lead fluoride precipitation but didn’t clarify the origin of the second exothermic event. Even if the crystallization of the overall glass matrix should be expected, further specific heat treatments and X-ray characterizations must be performed on this sample to fully describe the crystallization sequence. Finally, DSC measurements were performed on the bulk and powder glass samples 50PbEY and 60Pb to access the main crystallization mechanism which can be induced from nuclei at the surface (surface crystallization) or in the bulk (volume crystallization). When the crystallization process is induced from the surface, a large dependence of crystallization peak intensity and temperature is observed in function of the glass particle size. On the other hand, a preferential bulk volume crystallization can be identified by a similar crystallization behavior of the studied crystalline phase in the powder and bulk form. These results are presented in Figure 6. For sample 50PbEY, the very small shift of the onset of crystallization for both crystallization events is a strong indication of a preferential bulk nucleation of the corresponding crystalline phases, the first one being already attributed to PbF2(P04)3. In the case of composition 60Pb, the first crystallization peak due to cubic lead fluoride precipitation also suffers a very small temperature shift with particle size, indicating again a dominant bulk nucleation for this crystalline phase. On the other hand, a strong temperature shift can be observed for the high temperature crystallization event, suggesting that this unidentified crystalline phase is dominantly nucleated by the surface. Since a homogeneous bulk nucleation is required for control of the crystallite growth and distribution, these thermal results are promising and suggest that this glass composition is suitable for preparation of transparent glass-ceramics containing low phonon energy β-PbF2. However, this glass composition could not be doped with rare earth ions, which is a requirement for the desired applications but such doping may be achieved for lower rare earth concentrations or by varying the NaPO4/WO3 ratio under constant lead fluoride content in order to improve the glass stability against devitrification.

4. Conclusion

Er3+/Yb3+ codoped glass samples could be obtained in the ternary system (100-x)[0.6NaPO3-0.4WO3]-xPbF2 with x ranging from 0 to 50 and an undoped glass sample with x=60 could also be prepared. The most PbF2, concentrated samples (x=40, x=50 and x=60) were selected for thermal and crystallization characterizations. Thermal properties obtained by DSC for all samples pointed out a decrease of the glass transition temperature and glass thermal stability against devitrification related with the modifier behavior of lead fluoride. Heat-treatments performed near the crystallization peaks and X-ray diffraction measurements on the annealed samples indicated that the glass samples containing 40% and 50% of PbF2 preferentially precipitate lead fluorophosphate PbF2(P04)3 whereas the glass containing 60% of PbF2 presents a first crystallization event due to β-PbF2 nucleated homogeneously in the bulk and without other undesirable crystalline phase. These results suggest that this glass composition is promising for transparent glass-ceramics containing β-PbF2 nanocrystals for optical applications.

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