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# Glass-forming ability and ZrO<sub>2</sub> saturation limits in the magnesium aluminosilicate system

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## Abstract

Containerless melting was employed to evaluate the influence of increasing ZrO<sub>2</sub> additions on the glass-forming ability of peraluminous melts in the magnesium aluminosilicate system. The saturation limit of this oxide in glasses along the silica-cordierite join (SiO<sub>2</sub>–Mg<sub>2</sub>Al<sub>4</sub>Si<sub>5</sub>O<sub>18</sub>) is inversely proportional to the SiO<sub>2</sub> content; however, a limited solubility of ZrO<sub>2</sub> in melt-quenched silica exists (~2 mol%). Peraluminous samples (particularly MgO-free samples) exhibited a substantially higher capability of incorporating ZrO<sub>2</sub> without undergoing devitrification during cooling. TEM investigations revealed phase separation in all amorphous samples off the MgO = Al<sub>2</sub>O<sub>3</sub> line, with ZrO<sub>2</sub> preferentially segregating into a SiO<sub>2</sub>-depleted phase. These results identify the key role of Al<sub>2</sub>O<sub>3</sub> for the structural incorporation of ZrO<sub>2</sub> in silicate melts, opening up alternative pathways for the development of glasses and glass-ceramics respectively based on the homogeneous distribution of ZrO<sub>2</sub> or, in turn, on its exploitation to induce phase separation and crystal nucleation.

**Keywords:** glass solubility, ZrO<sub>2</sub> nucleation, phase separation, glass-forming ability, containerless melting

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## 1. Introduction

ZrO<sub>2</sub>-containing silicate melts remarkably constitute the common precursor for a variety of different functional materials. It is the case of electrofused refractory ceramics, whose long-lived chemical and thermal stability essentially relies on the complex assemblage of monoclinic ZrO<sub>2</sub> (m-ZrO<sub>2</sub>) and  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> crystallized from a parent melt during slow cooling [1]. As a widespread secondary component of glasses, ZrO<sub>2</sub> is moreover known to critically enhance their chemical durability [2,3], with a sensible impact on the corrosion mechanism of glasses for nuclear waste immobilization [4,5]. In glass-ceramics, ZrO<sub>2</sub> is additionally widely employed as a nucleating agent: during thermal annealing of the starting glass, it typically induces the early precipitation of nanosized tetragonal ZrO<sub>2</sub> crystals (t-ZrO<sub>2</sub>) [6–11], which act as heterogeneous nucleation centers for the subsequent controlled crystallization of the silicate matrix. Intensive research efforts have been more recently directed to the development of glass-ceramics with high crystalline ZrO<sub>2</sub> contents [12–16], to obtain rare-earth-free photoluminescence [17] or in the attempt of replicating the well-known transformation toughening typically observed in conventional ZrO<sub>2</sub> ceramics [18].

For all of the above-mentioned applications, a strict control over devitrification during cooling from the molten state must be achieved, which is particularly challenging at very high ZrO<sub>2</sub> content. Although geoscientists already developed reliable models to estimate the solubility of ZrO<sub>2</sub> in silicate melts as a function of temperature and composition [19–23], the studied compositions differed markedly from those of technologically relevant materials; more importantly, glass-forming ability was never directly evaluated. This study aims therefore at providing an enhanced systematic understanding of the quenchability of ZrO<sub>2</sub>-containing aluminosilicate glasses, to support the future compositional design of functional materials based on this metal oxide. To minimize heterogeneous surface nucleation of the silicate matrix and specifically examine the tendency of ZrO<sub>2</sub> to precipitate homogeneously

in the melts during quenching, we synthesized our samples by aerodynamic levitation coupled to laser heating (ADL) [24,25]. This containerless technique was indeed previously shown to enable the obtainment of exotic glasses with poor glass-forming ability (e.g. titanates, niobates and wolframates [26,27]). The results of our study complement the existing picture of the quaternary system  $\text{MgO-Al}_2\text{O}_3\text{-SiO}_2\text{-ZrO}_2$  and of its inherent tendency towards heterogeneity and phase separation, introducing alternative compositional approaches in the quest for superior glasses and glass-ceramics.

## 2. Materials and methods

Table 1. Nominal glass compositions analyzed on the peraluminous side of the  $\text{MgO-Al}_2\text{O}_3\text{-SiO}_2$  system, associated to all  $\text{ZrO}_2$  contents (e.g. 8 mol%  $\text{ZrO}_2$  + 92 mol% S55r1 in sample S55r1Zr8) tested to investigate their glass-forming ability. Devitrified compositions are marked by an \*.

| Sample name | Nominal composition of the base glass |                                |                       | $\text{ZrO}_2$ content(s) (mol%) |
|-------------|---------------------------------------|--------------------------------|-----------------------|----------------------------------|
|             | MgO (mol%)                            | $\text{Al}_2\text{O}_3$ (mol%) | $\text{SiO}_2$ (mol%) |                                  |
| S100        | 0                                     | 0                              | 100                   | 0, 0.75, 1.5, 2*, 4*             |
| S85r1       | 7.5                                   | 7.5                            | 85                    | 0, 2, 4*, 8*                     |
| S85r0       | 0                                     | 15                             | 85                    | 0, 4                             |
| S75r1       | 12.5                                  | 12.5                           | 75                    | 0, 2, 4, 8*                      |
| S75r0.4     | 7                                     | 18                             | 75                    | 8                                |
| S75r0       | 0                                     | 25                             | 75                    | 0, 4, 8                          |
| S55r1       | 22.5                                  | 22.5                           | 55                    | 0, 4, 6, 8, 10, 15*, 20*         |
| S55r0.5     | 14                                    | 31                             | 55                    | 15*                              |
| S55r0.2     | 7                                     | 38                             | 55                    | 15                               |
| S55r0       | 0                                     | 45                             | 55                    | 0, 8, 15                         |

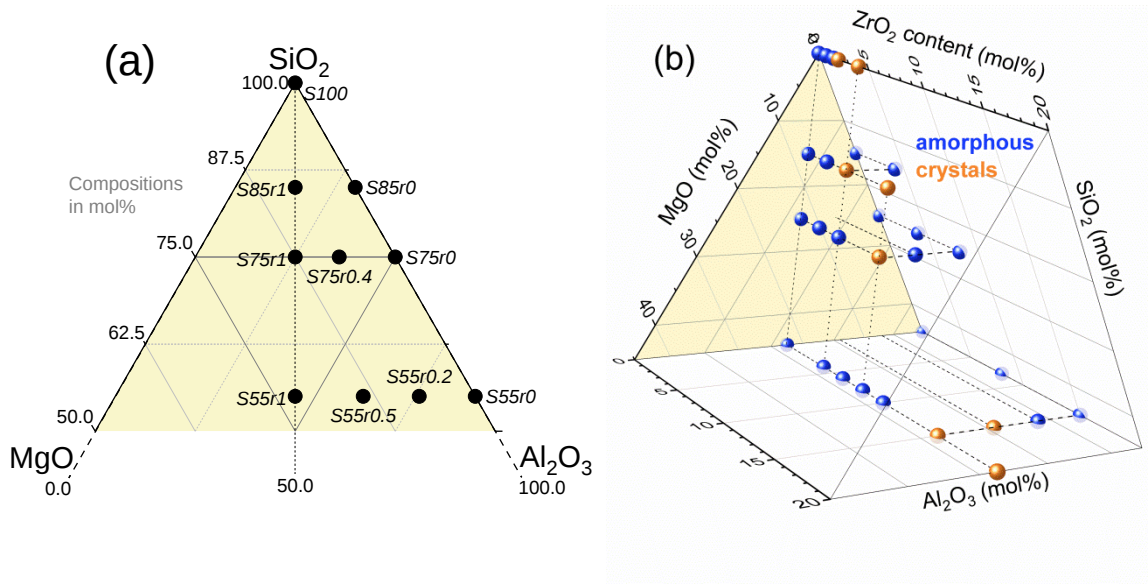


Figure 1. a) Upper section of the MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> ternary diagram, reporting the base glass compositions analyzed within this work. b) Summary of all samples synthesized within this work (see also Table 1): partially devitrified materials (as concomitantly determined by XRD and Raman) are identified by orange spheres, while fully amorphous samples are shown in blue (mind that all amorphous samples off the line MgO = Al<sub>2</sub>O<sub>3</sub> were investigated by TEM and found to exhibit phase separation; see Fig. 5 and related discussion).

## 2.1 Synthesis of the samples

Our study focused on the peraluminous side of the MAS system (see Tab. 1 and Fig. 1); attempts to extend the examination towards strongly subaluminous compositions invariably yielded opaque or devitrified samples. Base compositions S100, S85r1, S75r1 and S55r1 located on the silica-cordierite join, along which the MgO/Al<sub>2</sub>O<sub>3</sub> molar ratio equals 1; other samples were then defined by substituting some (or all) their MgO with Al<sub>2</sub>O<sub>3</sub>. The chosen nomenclature reflected these compositional features, e.g. S55r0.2 for a sample with 55 mol% SiO<sub>2</sub> and MgO/Al<sub>2</sub>O<sub>3</sub> = 0.2. An additional suffix manifested the addition of ZrO<sub>2</sub> (S55r0.2Zr15 for 15 mol% ZrO<sub>2</sub> added to 85 mol% S55r0.2 glass). All samples were synthesized using laboratory-grade raw materials: MgO (99.5%, Strem Chemicals), Al<sub>2</sub>O<sub>3</sub>

(99.999%, Strem Chemicals), SiO<sub>2</sub> (99.9%, Chempur) and ZrO<sub>2</sub> (99%, Koch-Light Laboratories). The final melt-quenching stage took place invariably by ADL in an Ar gas flow, using a top and a bottom CO<sub>2</sub> lasers (wavelength: 10.6 μm; maximum power: 250 W) to heat up the samples and two pyrometers to estimate their temperature; the average quenching rate was computed at ~250 K s<sup>-1</sup> (see Figs. S1 and S2 in SI section). The synthesis protocol was however adjusted to each composition:

- in the case of the highly viscous S100 series, batches of 5 g (SiO<sub>2</sub>+ZrO<sub>2</sub>) were mixed for 3 h in an agate ball mill to achieve thorough homogenization. After this, the powders were pressed into pellets, from which small chunks (10-20 mg) were detached and melted by ADL at 1900-2100 °C for approximately 5 s; quenching was simply achieved by shutting off the lasers. After crushing and grinding the glassy (or devitrified) beads in a hand mortar, the procedure was repeated twice.
- given their lower melting temperature, S85r1, S75r1 and S55r1 were first produced as ZrO<sub>2</sub>-free glasses by conventional furnace melting in Pt-Rh20 crucibles: after mechanical mixing for 3 h, batches of 4 g were heated at 1650 °C for 2 h, quenched, crushed and remelted to ensure homogeneity. After this, ZrO<sub>2</sub> was added to approximately 2 g of the obtained glasses and the double furnace melting at 1650 °C was repeated in the same way. Finally, all samples were melted once by ADL.
- the peraluminous samples, exhibiting high liquidus temperatures but comparatively low viscosity, were melted only once by ADL at 1900-2100 °C, after carefully mixing the starting powders in an agate hand mortar and pressing pellets of 1 g.

To compare the Raman features of ZrO<sub>2</sub> crystals formed during melt-quenching or secondary heat treatments, three glassy samples were additionally annealed in the furnace at selected temperatures: 15 min at 1000 °C for S75r1Zr4, 15 min at 900 °C for S55r1Zr8 and 1 h at 1000 °C for S55r0Zr15. We estimated these temperatures based on previous works

[28,29] and own DSC and high temperature XRD measurements (see Figs. S4 and S5 in the SI section). The three samples are labeled with an additional *-ht* suffix.

## 2.2 Raman Spectroscopy

The as-prepared glassy beads were measured on a Renishaw InVia Qontor spectrometer equipped with a green laser (514 nm) operating at 50 mW (the low-wavenumber cutoff of the edge filter is at  $\sim 100\text{ cm}^{-1}$ ). After focusing 20-40  $\mu\text{m}$  below the surface, the spectra were collected in the range  $80\text{-}2000\text{ cm}^{-1}$  using a 1800 lines/mm holographic grating, with 15 s acquisition time and 6 repetitions. Laser power and acquisition time were occasionally lowered to avoid detector saturation in the case of highly crystalline materials. After applying a simple baseline, the spectra were normalized to their maximum intensity to facilitate comparison.

## 2.3 X-ray Diffraction (XRD)

XRD characterization was performed in Bragg-Brentano geometry using a D8 Advance Bruker laboratory diffractometer, equipped with a Cu X-ray tube and a LynxEye XE line detector. Single glassy beads were crushed and grinded in an agate hand mortar and the resulting powders were dispersed on low-background flat Si sample holders with the help of some ethanol drops. The measurements were run at 30 kV – 40 mA in the range  $15\text{-}80^\circ 2\theta$ , with increments of  $0.025^\circ 2\theta$  and 0.7 s for each step. In-situ high-temperature XRD measurements (Figure S3 of the SI section) were performed to identify suitable temperatures for the heat treatments described in Section 2.1, using Anton Paar's HTK1200N heating chamber. The powdered glass samples were deposited on a platinum disk to avoid any reaction with the underlying alumina crucible and data were collected every  $50^\circ\text{C}$  between  $600^\circ\text{C}$  and  $1200^\circ\text{C}$  with Vantec-1 line detector.

## 2.4 Transmission electron microscopy (TEM)

Single beads were crushed and grinded in an agate hand mortar and the resulting powder was then dispersed in ethanol. A small drop of the suspension was loaded on a TEM copper grid layered by an amorphous holey carbon film and dried. Imaging of the samples and elemental analyses were performed with a Philips CM20 TEM operated at 200 kV and fitted with an EDAX EDS system.

## 3. Results

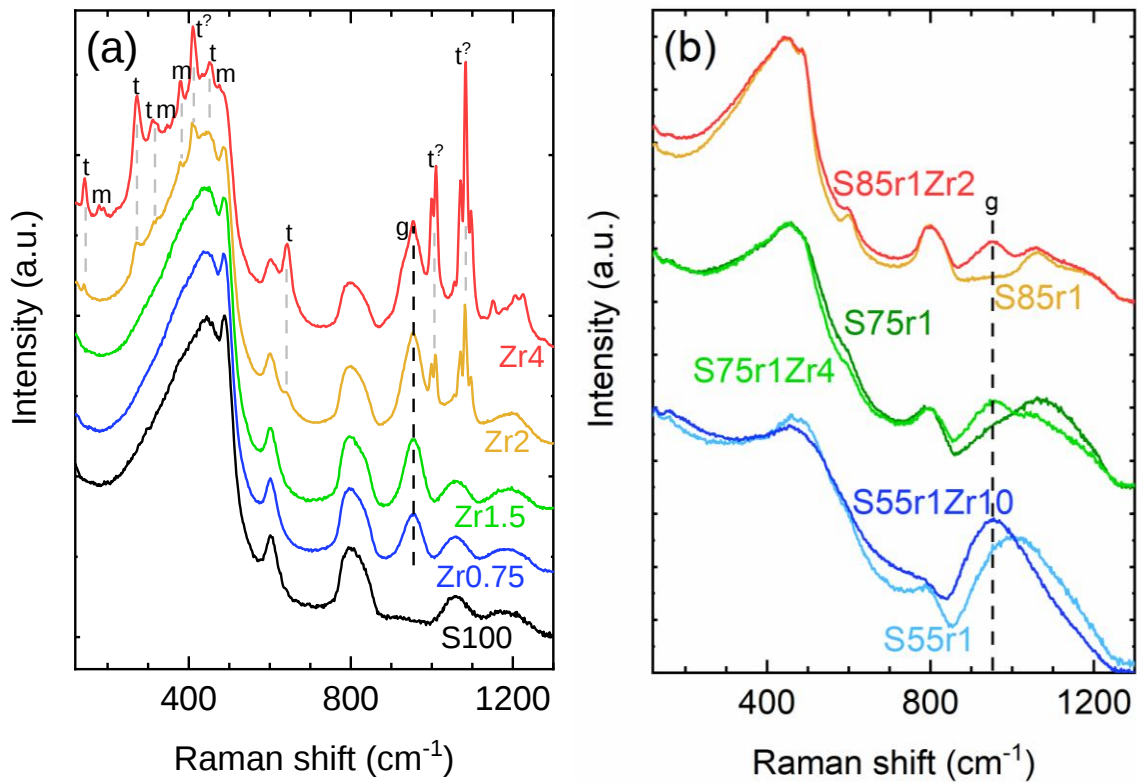


Figure 2. Raman spectra collected from: a) samples of base composition S100, doped with increasing amounts of  $\text{ZrO}_2$  (0-4 mol%); b) glasses of base composition S85r1, S75r1 and S55r1 with and without  $\text{ZrO}_2$ , in the maximum investigated addition still yielding an

amorphous material. In both cases, *g* marks the position of the most intense Raman feature associated with the incorporation of  $\text{Zr}^{4+}$  in the amorphous (alumino)silicate network; other labels: *t* for t-ZrO<sub>2</sub> [30], *t*<sup>2</sup> for bands clearly associated with t-ZrO<sub>2</sub> but absent in the literature (see discussion in the text), *m* for m-ZrO<sub>2</sub> [30].

Raman measurements were used to verify the successful incorporation of ZrO<sub>2</sub> in the amorphous structure of the synthesized (alumino)silicate glasses. As expected, the spectrum of undoped S100 (Fig. 2-a) exhibited the typical vibrational features of pure SiO<sub>2</sub>, i.e. broad asymmetric bands at 450 and 800 cm<sup>-1</sup> associated with the vibrations of Si-O-Si linkages, sharper “defect” lines at 490 and 600 cm<sup>-1</sup> stemming from the breathing modes of four- and three-membered tetrahedral rings and, above 1000 cm<sup>-1</sup>, the envelope for the stretching of bridging oxygens about SiO<sub>4</sub> tetrahedra [31–35]. In turn, an additional relatively broad band was observed in the spectra of ZrO<sub>2</sub>-doped S100 samples, centered at ~950 cm<sup>-1</sup> and growing in intensity with increasing ZrO<sub>2</sub> content; a similar vibrational feature has been previously observed in SiO<sub>2</sub>-ZrO<sub>2</sub> gels [36], ZrO<sub>2</sub>-bearing borosilicate [37,38] and aluminosilicate glasses and assigned to the stretching of bridging oxygens about SiO<sub>4</sub> Q<sup>3</sup> units linked to a Zr polyhedron [39]. S100Zr2 and S100Zr4 could not be fully vitrified and accordingly display several supplementary sharp bands stemming from crystalline material (see treatment below).

Similar considerations can be applied to the Raman spectra of S85r1, S75r1 and S55r1 (Fig. 2-b): their undoped spectra manifested a consistent broadening at low wavenumbers and a gradual downshift at high wavenumbers with respect to S100, due to the progressive Al<sub>2</sub>O<sub>3</sub> and MgO enrichment of the glass. These observations agree with previous Raman investigations in the MAS system [40]. Again, the addition of ZrO<sub>2</sub> brought about the appearance of a supplementary vibrational feature at ~950 cm<sup>-1</sup>, which can be therefore univocally associated with the incorporation of this oxide in the amorphous (alumino)silicate

structure. The spectrum of the Al-rich samples (especially S55r1) appeared moreover to be more profoundly impacted by the  $\text{ZrO}_2$  doping, with a further smearing out of the original silicate bands and a clear downshift of the high-wavenumber vibrational envelope. Note that the shown spectra correspond to the highest experimented  $\text{ZrO}_2$  contents which yielded fully amorphous samples along the silica-cordierite join (see Figures S6, S7 and S8 of the SI section for the spectra of all synthesized samples).

The only crystalline phase contained in samples located slightly above the  $\text{ZrO}_2$  saturation limit (Fig. 1-b) was invariably t- $\text{ZrO}_2$ , as confirmed by the comparison of the observed Raman bands (at roughly 145, 270, 320, 450, 600 and 650  $\text{cm}^{-1}$  in Figs. 2-a and 3) with previous references [30]. Despite the lack of literature correspondence, the sharp features observed at 410  $\text{cm}^{-1}$  and above 1000  $\text{cm}^{-1}$  (see Discussion) were clearly related to the same phase. This whole set of Raman bands appeared also after annealing above  $T_g$  samples that were initially obtained as glasses, as inferable from the spectra of S75r1Zr4ht and S55r1Zr8ht. At higher doping levels, the additional formation of m- $\text{ZrO}_2$  took place during quenching, such as in S55r1Zr20 and S100Zr4. These results were corroborated by parallel XRD measurements (Fig. 3-b).

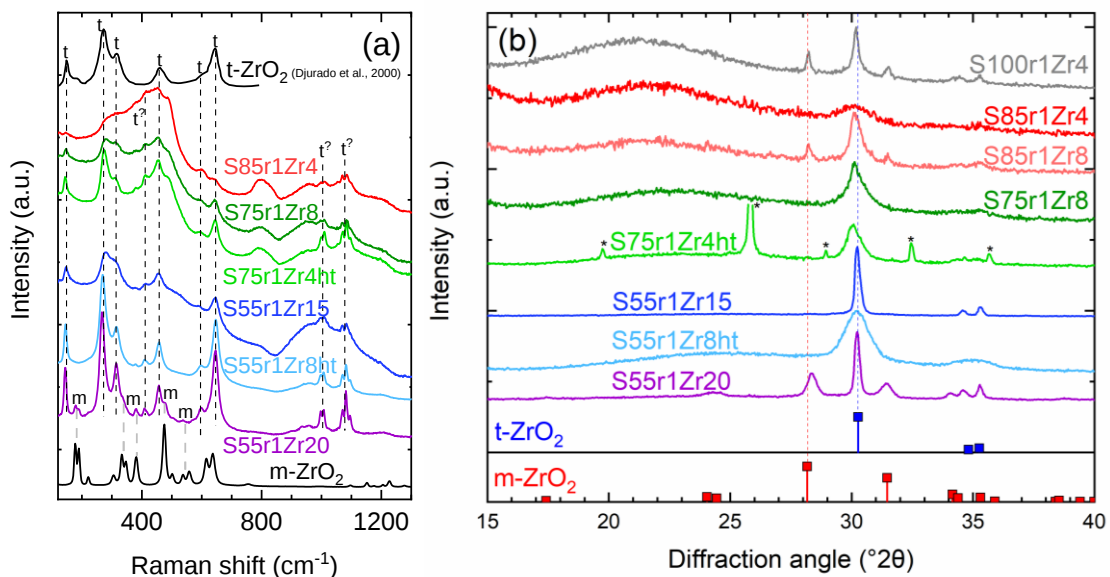


Figure 3. a) Raman spectra collected from samples of S85r1, S75r1 and S55r1 base composition, devitrified during cooling from the melt or after controlled secondary heat treatments (-*ht* suffix); the spectrum of the employed m-ZrO<sub>2</sub> raw material is provided for comparison. Labels: *t* for t-ZrO<sub>2</sub> [30], *t*<sup>2</sup> for bands clearly associated with t-ZrO<sub>2</sub> but absent in the literature (see discussion in the text), *m* for m-ZrO<sub>2</sub> [30]. b) XRD measurements performed on several as-quenched and heat-treated samples in comparison to the PDF cards of m-ZrO<sub>2</sub> [41] [ICDD 00-037-1484] and t-ZrO<sub>2</sub> [42] [ICDD 00-050-1089].

Interestingly, the capability of peraluminous (and MgO-free) samples to incorporate ZrO<sub>2</sub> in a fully amorphous structure was found to be significantly higher than that of their counterparts along the silica-cordierite join. While S85r1Zr4 contained t-ZrO<sub>2</sub>, S85r0Zr4 was completely amorphous (Fig. 4); at lower SiO<sub>2</sub> content, S75r1Zr8 located above the glass saturation limit for ZrO<sub>2</sub>, while S75r0.4Zr8 and S75r0Zr8 showed no visible sign of devitrification. The Al<sub>2</sub>O<sub>3</sub> excess particularly emerged as crucial when examining the S55 series: S55r1Zr15 and S55r0.5Zr15 devitrified during cooling, whereas the Al<sub>2</sub>O<sub>3</sub>-richer samples S55r0.2Zr15 and S55r0Zr15 were totally amorphous. The Raman spectra underwent again a gradual broadening of their features (including the characteristic band at ~950 cm<sup>-1</sup>) as the SiO<sub>2</sub> content of the glasses reduced.

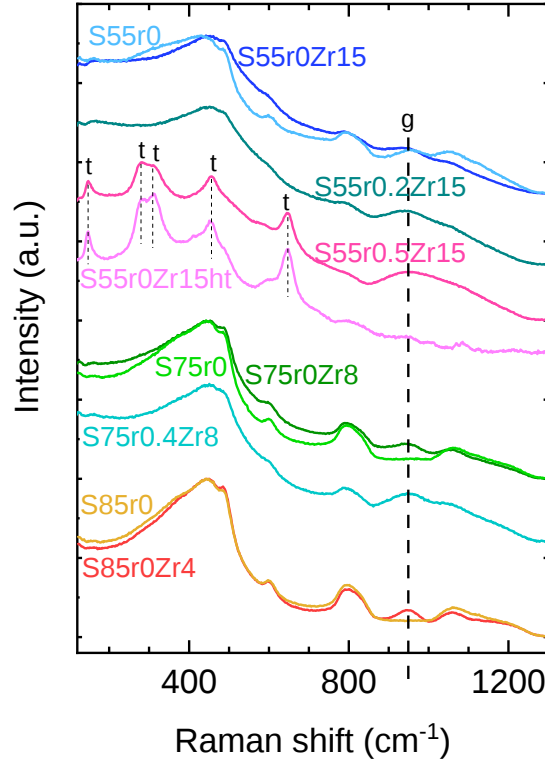


Figure 4. Raman spectra of various samples quenched on the peraluminous side of the MgO- $\text{Al}_2\text{O}_3$ - $\text{SiO}_2$  ternary system, with and without  $\text{ZrO}_2$  doping; *g* signals the position of the most intense Raman feature associated with the incorporation of  $\text{Zr}^{4+}$  in the amorphous (alumino)silicate network, *t* for t- $\text{ZrO}_2$  [30].

All amorphous samples with an excess of  $\text{Al}_2\text{O}_3$  over MgO were analyzed by TEM to investigate their nanostructure, since base compositions S75r0 and especially S55r0 and S55r0.2 exhibited a noticeable opalescence with or without the addition of  $\text{ZrO}_2$  (Fig. S3 of the SI section). Bright-field micrographs (Fig. 5) revealed the emergence of pervasive amorphous phase separation in all these peraluminous samples, embodied by the intergrowth of a high contrast (and therefore  $\text{ZrO}_2$ -rich) phase and a low-contrast one. Some of the materials (e.g. S85r0Zr4, S55r0.2Zr15, S55r0Zr15) exhibited a stronger analogy to

nanostructures typically assigned to a nucleation-growth mechanism, whereas the others matched more closely with the result of a spinodal decomposition or exhibited an intermediate character [43]. In sample S85r0 particularly, phase separation was only incipient and displayed diffused interfaces, making its recognition rather challenging. The average size of phase-separated domains (Fig. 5-g) was estimated in all samples by measuring the diameter of pseudo-spherical particles (or the smallest dimension of irregularly shaped ones, assuming them to originate from coalescence) located in the thinnest electron-transparent regions (to avoid artifacts due to object overlap). The resulting values enabled to explain the different optical behaviors of the samples, as only the biggest sizes induced a noticeable opalescence, in agreement with the available models [44]. The following relations could be additionally extracted: (i) in absence of  $\text{ZrO}_2$ , the phase separation increased in size with an increasing Al/Si ratio; (ii) at a given Al/Si ratio, the addition of  $\text{ZrO}_2$  brought about a substantial increment in size and contrast of the demixed domains; (iii) substitution of  $\text{Al}_2\text{O}_3$  by MgO (at given  $\text{ZrO}_2$  and  $\text{SiO}_2$  contents) led to the opposite effect. Thanks to the particularly big size of phase separation in sample S55r0Zr15, EDX enabled also a reliable estimation of the composition of its demixed regions: the low-contrast droplets yielded an average of 63(6) mol%  $\text{SiO}_2$ , 25(3) mol%  $\text{Al}_2\text{O}_3$  and 12(1) mol%  $\text{ZrO}_2$ , while the result for the enclosing high-contrast matrix was 27(2) mol%  $\text{SiO}_2$ , 50(1) mol%  $\text{Al}_2\text{O}_3$  and 23(2) mol%  $\text{ZrO}_2$ .

We additionally characterized by TEM sample S85r1Zr4, which devitrified during cooling and was macroscopically transparent (see Figs. S3 and S10 of the SI section). It contained a population of crystalline nuclei with an average size between 5 and 10 nm in an otherwise homogeneous glass matrix. Selected area electron diffraction (SAED) patterns exhibited continuous diffuse rings and provided  $d$ -spacings assignable to the structure of  $t$ - $\text{ZrO}_2$  (the two examples highlighted on Fig. S10 yielded 1.8(1) Å and 3.0(1) Å, respectively

corresponding to the (011) and (112) planes of t-ZrO<sub>2</sub> [42]), in agreement with Raman and XRD results.

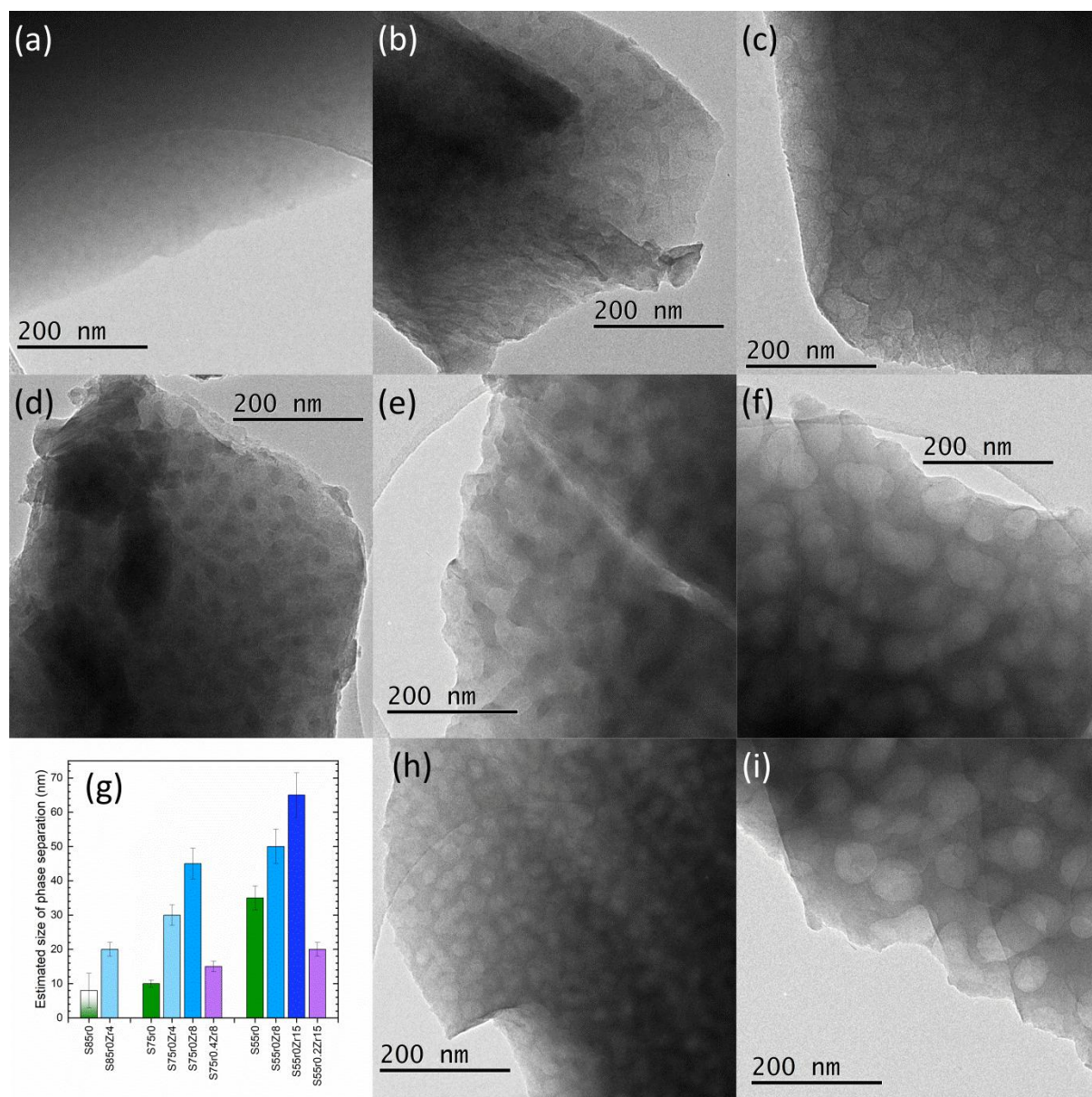


Figure 5. TEM bright-field micrographs of phase-separated samples: a) S85r0, b) S75r0, c) S55r0, d) S85r0Zr4, e) S75r0Zr8, f) S55r0Zr8, h) S55r0.2Zr15 and i) S55r0Zr15. g) Average size of the phase-separated regions in each sample, as estimated from TEM micrographs (see also Fig. S10 in the SI section).

#### 4. Discussion

Containerless melting enabled us to screen a wide range of compositions and  $\text{ZrO}_2$  doping levels in silicate melts, overcoming the temperature limitations of conventional melt-quenching and relating glass-forming ability purely to the tendency of  $\text{ZrO}_2$  to nucleate homogeneously upon cooling. Our approach was mainly concerned with the identification of the  $\text{ZrO}_2$  saturation limit in glasses quenched from above their liquidus temperatures; therefore, its results can be only limitedly compared with those of geoscientists investigating the subliquidus melt solubility of zircon as a function of temperature and composition [19–23]. However, our study confirms the key relevance of  $\text{SiO}_2$  content and overall structural polymerization for the incorporation of  $\text{Zr}^{4+}$  in silicate glasses: the saturation threshold of this oxide increased steadily along the silica-cordierite join and was shown to be even higher at peraluminous compositions. Clearly, this must be due to the difficulty of a fully polymerized  $[\text{SiO}_4]$  tetrahedral network to accommodate high amounts of  $\text{Zr}^{4+}$ , which has been shown to assume preferentially a 6- or 7-fold oxygen coordination in  $\text{SiO}_2$ - $\text{ZrO}_2$  xerogels and in aluminosilicate glasses, sharing edges with adjacent (Si,Al)-polyhedra [6,9,39,45].

In a subaluminous soda lime glass,  $\text{ZrO}_2$  oversaturation was previously correlated to a gradual shift from 6- to 8-fold oxygen coordination around  $\text{Zr}^{4+}$  ions, which would therefore prove less efficient than  $\text{Al}^{3+}$  ions (invariably 4-coordinated even at high  $\text{ZrO}_2$  content) in the competition for charge-balancing modifier cations [39]. In MAS glasses, however, 5- and 6-fold-coordinated  $\text{Al}^{3+}$  species have been shown to be predominant even in the absence of  $\text{ZrO}_2$ , especially at peraluminous compositions [40]: the presence of these highly coordinated, modifier-like  $\text{Al}^{3+}$  species (and the resulting lower network rigidity) may therefore represent a key factor for the solubility of  $\text{ZrO}_2$  in the glasses studied within this work. In fact, Raman spectra of our  $\text{Al}_2\text{O}_3$ -rich samples highlighted more extensive structural responses to the addition of  $\text{ZrO}_2$  (the noted broadening and shift of Raman features); on the contrary, the spectrum of  $\text{SiO}_2$  appeared barely affected by the dopant if not through the appearance of the

additional characteristic band at  $\sim 950\text{ cm}^{-1}$ , previously associated to occurrence of linkages between  $Q^3$  units and  $Zr^{4+}$  oxygen polyhedra [39].

The inferred favorability of an  $Al_2O_3$ -rich environment for the amorphous incorporation of  $Zr^{4+}$  is reinforced by our TEM investigations. Phase separation could be observed already in samples belonging to the binary  $Al_2O_3$ - $SiO_2$  system, in which a metastable sub-liquidus immiscibility field is known from literature [46]. These materials were likely to demix during quenching, despite the relatively fast cooling rate of our ADL system (Fig. S2 of the SI section); the increment in size of the demixed regions with an increasing Al/Si ratio may therefore be associated to the expectably higher liquidus temperature and lower viscosity of  $Al_2O_3$ -richer melts, fostering nucleation and growth. As for  $Zr^{4+}$ , it was found to preferentially segregate in phase-separated regions with the highest Al/Si ratio, further increasing their size. This is indeed not surprising if one considers that a glass-forming region has been previously determined by plasma-spraying or melt-extraction at intermediate compositions of the binary  $Al_2O_3$ - $ZrO_2$  system [47–49]. On the contrary, an addition of MgO appeared to reduce the extent of phase separation, in analogy to the effect of alkali oxides such as  $Na_2O$  previously reported in similarly immiscible systems [50]. Other authors reported unavoidable nanoscale chemical heterogeneity (below the resolution of our TEM) even in a melt-quenched glass with virtually the same stoichiometry as our S75r1Zr4 sample [28], in which MgO equals  $Al_2O_3$ . The tendency towards heterogeneity and phase separation may therefore be inherent to the whole MgO- $Al_2O_3$ - $SiO_2$  system, especially if  $ZrO_2$  is added to compositions locating between the above-mentioned  $Al_2O_3$ - $SiO_2$  metastable immiscibility field [46] and, on the other side, a stable liquid immiscibility region in the MgO- $SiO_2$  binary system [51]. The heterogeneous distribution of  $ZrO_2$  at the nanoscale was also related to the ability of these materials to nucleate t- $ZrO_2$  crystals during secondary annealing and yield technologically relevant glass-ceramics [9,28,52]. Overall, our results

complement the available literature and provide a solid fundamental starting point for the future compositional design of homogeneous glasses containing  $\text{ZrO}_2$  or, in turn, for the exploitation of incipient heterogeneity and phase separation in the quest for novel transparent glass-ceramic materials, as recently demonstrated in immiscible zinc gallogermanate glasses [50].

Concerning now the  $\text{ZrO}_2$  crystals in the devitrified samples, our observations are in line with the works of previous authors:  $t\text{-ZrO}_2$  was already shown to precipitate preferentially from  $\text{SiO}_2\text{-ZrO}_2$  plasma-sprayed melts [47] and the occurrence of this metastable polymorph was explained in terms of an energy minimization due to the high surface energy of the nanocrystals and/or associated to the mechanical constraints exerted by the surrounding glassy phase [30,47,53]. A chemical stabilization by  $\text{Al}_2\text{O}_3$ - and/or  $\text{MgO}$ -doping was instead previously speculated in the MAS system [54]: this might be at the origin of the substantial broadening of the  $t\text{-ZrO}_2$  Raman bands in our MAS samples (Fig. 3), especially if compared with the spectra of S100Zr2 and S100Zr4 (Fig. 2). In turn, the additional bands observed in our spectra at  $410\text{ cm}^{-1}$  and above  $1000\text{ cm}^{-1}$ , though absent in the literature references for  $t\text{-ZrO}_2$  [30], were clearly related to the formation of this phase, the only one detectable by XRD. They may be tentatively assigned to the presence of Si-O-Zr linkages in the nanocrystals, given the similarity of their positions with those of the main bands of zircon  $\text{ZrSiO}_4$  (see for example [RRUFF ID: R050034] [55]), their peculiar sharpness in S100 samples and, conversely, their absence in  $\text{Al}_2\text{O}_3$ -rich samples such as S55r0Zr15ht and S55r0.5Zr15.

Crystals formed during melt devitrification or secondary glass annealing appeared virtually identical in our Raman spectra; similarly, the TEM micrographs of S85r1Zr4 (Fig. S10), devitrified during cooling, closely resembled the homogeneously distributed nanocrystals typically encountered in glass-ceramics. It is therefore possible to conclude that

ZrO<sub>2</sub> crystallization must have taken place at relatively deep undercooling during quenching, probably driven by the temperature-dependent oversaturation of the melt and by the gradual slow-down of the quenching rate in our ADL system (Fig. S2 in the SI section). Such a crystallization mechanism may indeed open up more direct pathways to obtain transparent nanocrystallized materials by controlled cooling from the melt, bypassing the vitrification stage which is presently inevitable in conventional glass-ceramic production [56]. On the contrary, the adoption of a different levitation gas with higher thermal conductivity (such as O<sub>2</sub>) and the production of smaller glass beads could have probably enabled an even higher ZrO<sub>2</sub> incorporation than documented in this study, by increasing the quenching rate; nevertheless, the identified compositional trends would still be valid. Regarding the appearance of m-ZrO<sub>2</sub> at higher degrees of supersaturation, it is to be attributed to the bigger ZrO<sub>2</sub> crystallite size or, possibly, to the presence of unmelted residues.

## 5. Conclusion

Our study has shed new light on the chemical constraints influencing the saturation threshold of ZrO<sub>2</sub> in aluminosilicate melts, paving the way for a smarter compositional design of technical glasses and glass-ceramics based on this oxide, e.g. by controlled cooling from the melt, exploiting phase separation or by conventional glass-ceramic processing. SiO<sub>2</sub> content and overall polymerization degree were found to be the main factors influencing the successful incorporation of Zr<sup>4+</sup> in an amorphous network, with a crucial role played by the structural versatility of Al<sup>3+</sup>. Concurrently, the chemical heterogeneity of as-quenched materials emerged once more as a preeminent feature of MAS glasses doped with ZrO<sub>2</sub>, evoking alternative compositional approaches for the development of novel functional materials.

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## Figure Captions

Figure 1. a) Upper section of the MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> ternary diagram, reporting the base glass compositions analyzed within this work. b) Summary of all samples synthesized within this work (see also Table 1): partially devitrified materials (as concomitantly determined by XRD and Raman) are identified by orange spheres, while fully amorphous samples are shown in blue (mind that all amorphous samples off the line MgO = Al<sub>2</sub>O<sub>3</sub> were investigated by TEM and found to exhibit phase separation; see Fig. 5 and related discussion).

Figure 2. Raman spectra collected from: a) samples of base composition S100, doped with increasing amounts of ZrO<sub>2</sub> (0-4 mol%); b) glasses of base composition S85r1, S75r1 and S55r1 with and without ZrO<sub>2</sub>, in the maximum investigated addition still yielding an amorphous material. In both cases, *g* marks the position of the most intense Raman feature associated with the incorporation of Zr<sup>4+</sup> in the amorphous (alumino)silicate network; other labels: *t* for t-ZrO<sub>2</sub> [30], *t*<sup>2</sup> for bands clearly associated with t-ZrO<sub>2</sub> but absent in the literature (see discussion in the text), *m* for m-ZrO<sub>2</sub> [30].

Figure 3. a) Raman spectra collected from samples of S85r1, S75r1 and S55r1 base composition, devitrified during cooling from the melt or after controlled secondary heat treatments (-*ht* suffix); the spectrum of the employed m-ZrO<sub>2</sub> raw material is provided for comparison. Labels: *t* for t-ZrO<sub>2</sub> [30], *t*<sup>2</sup> for bands clearly associated with t-ZrO<sub>2</sub> but absent in the literature (see discussion in the text), *m* for m-ZrO<sub>2</sub> [30]. b) XRD measurements performed on several as-quenched and heat-treated samples in comparison to the PDF cards of m-ZrO<sub>2</sub> [41] [ICDD 00-037-1484] and t-ZrO<sub>2</sub> [42] [ICDD 00-050-1089].

Figure 4. Raman spectra of various samples quenched on the peraluminous side of the MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> ternary system, with and without ZrO<sub>2</sub> doping; *g* signals the position of the most intense Raman feature associated with the incorporation of Zr<sup>4+</sup> in the amorphous (alumino)silicate network, *t* for t-ZrO<sub>2</sub> [30].

Figure 5. TEM bright-field micrographs of phase-separated samples: a) S85r0, b) S75r0, c) S55r0, d) S85r0Zr4, e) S75r0Zr8, f) S55r0Zr8, h) S55r0.2Zr15 and i) S55r0Zr15. g) Average size of the phase-separated regions in each sample, as estimated from TEM micrographs (see also Fig. S10 in the SI section).