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Thio and selenosilicates, sulfide and selenide counterparts of silicates: similarities and differences

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Abstract. Thiosilicate and selenosilicate glasses are the sulfide and selenide counterparts of the well-known silicate glasses. This paper reviews the main investigations carried out to shed light on the structural, thermal and electrical characteristics of these glasses, showing their particularities compared to their oxide counterparts, such as a more complex structure with the presence of both, corner- and edge-sharing tetrahedra and much higher ion conductivity due to the larger polarizability of the chalcogen and the more covalent nature of the bonds, but also, their similarities with the presence of both, bridging and non-bridging S or Se or the existence of a mixed alkali effect when two mobile Li⁺ and Na⁺ cations coexist in the glassy network. Structural consequences of competition between SiS₂ and another former, GeS₂, PS₅/₂ or BS₃/₂ are also discussed.

Keywords. Thiosilicate, Selenosilicate, Edge-sharing tetrahedron, Mixed alkali effect, Glass former competition, Glass structure, Solid electrolyte.

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1. Introduction

In the mid-sixties, considerable research efforts were devoted to the study of new materials with potential interest as solid electrolytes. At the time, focus was put on crystalline compounds. Very remarkable compounds were found such as sodium β-alumina, RbAg₄I₅, NASICON (Sodium Superionic Conductor). Not much interest was reserved for glasses since the conductivity of ion-conducting glasses, the only known ones being oxides, was rather poor. The situation changed in the late seventies with the publication of the “Weak Electrolyte Theory” (WET) [Ravaine and Souquet, 1977].

In this model, the glass is treated as a weak electrolyte with the glass former compound, e.g. SiO₂, being the solvent and the modifier compound, e.g. Na₂O, being the solute, such assumption being supported by the fact that glasses have very low dielectric constants (εr ~ 5–10 in oxide glasses). The authors have established experimentally that the electrical conductivity of xNa₂O–(1–x)SiO₂ glasses varied as the square root of their thermodynamic activity. And, with the assumption of a weak dissociation, so does the concentration of Na⁺ ions resulting from the dissociation of the modifier compound (Na₂O ↔ Na⁺ + NaO⁻). Therefore, the conductivity varies as the concentration of these “free” ions. The authors proposed that the only charge carriers free to move throughout the glassy matrix in the presence of an electric field were those released as a result of the solute dissociation. A larger solute dissociation
would then result in a larger number of free charge carriers, and subsequently in a larger conductivity.

According to the WET, the ionic conductivity in a glass is then closely related to the dielectric permittivity of the medium and will increase with increase in this parameter. Therefore, the replacement of oxygen by a more polarizable atom, such as S, or Se, was supposed to increase the conductivity. The first Na⁺-conducting thiogermanate glasses were then synthesized [Barrau et al., 1978]. In the following years, owing to the success of the WET, ion-conducting glasses based upon other sulfide networks such as thioborate [Levasseur et al., 1981], thiophosphate [Mercier et al., 1981], thioarseniate [Visco et al., 1985] and thiosilicate networks [Akridge, 1984, Pradel and Ribes, 1986, Sahami et al., 1985] were investigated.

It was the first time that these families, sulfide counterparts of the very well-known silicate, germanate, borate and phosphate glasses were elaborated. At the time, simple glasses comprising a network and a modifier were prepared. Later, more complex systems were elaborated in order to investigate famous effects, largely debated in the oxide family, such as the mixed alkali effect (MAE), the mixed glass former effect (MFE). The motivation of these investigations being the search for fast ion conductors, literature reports mainly on glass systems comprising Li⁺ and Na⁺ as mobile cations.

In this paper, we will only focus on thiosilicate and selenosilicate glasses, and report their preparation, structural, thermal and electrical characterization, showing their particularities compared to their oxide counterparts due to the larger polarizability of the chalcogen and the more covalent nature of the bonds but also, their similarities with the existence of both, bridging and non-bridging anions or the non-linear variation of properties when two mobile cations coexist in the glassy network.

2. Elaboration of thio and selenosilicate glasses

Like their oxide counterparts, thiosilicate and selenosilicate glasses can be prepared as bulk materials, powders or thin films. On the other hand, and in contrast to oxides, their preparation requires an oxygen-free atmosphere. After preparation, the glasses must be handled in moisture-free glove box since they are highly hygroscopic.

The usual way to prepare the glasses is the direct reaction of the constituting elements (for the former compounds, Si and S or Se) or the reaction of the constituting modifier compounds (e.g. Li₂S), former compound (e.g. SiS₂, GeS₂…) and eventually “dopants” (e.g. LiI) mixed in stoichiometric quantities and placed in quartz tubes sealed under vacuum. The mixture is then melted at temperatures usually comprised between 1000 K and 1350 K, lower than the required temperatures for silicate glasses (\(T_f\) SiO₂ = 1983 K; \(T_f\) SiS₂ = 1373 K; \(T_f\) SiSe₂ = 1243 K). Depending upon the ease of vitrification, the tube is quenched in air or in various liquids that increase the heat dissipation (water, salted water, liquid nitrogen). When preparing alkali-conducting chalcogenide glasses, vitreous carbon crucibles are often inserted in the tube in order to avoid contact of the reacting mixture with the quartz tube and thus contamination of the glass by silica [Levasseur et al., 1981, Pradel and Ribes, 1986, Robinel et al., 1983]. The vitreous carbon crucible can be replaced by a layer of carbon obtained by pyrolysis of an organic compound such as acetone [Kennedy, 1989, Zhang and Kennedy, 1990].

Sulfur and to a lesser extend, selenium have high vapor pressures. A slow heating ramp of ∼6 K/hour (in presence of S or Se) and ∼30 K/hour (in presence of already prepared modifier, former compounds…) at the beginning of the reaction process is therefore required in order to avoid any risk of explosion of the quartz tube. The heat treatment can include steps with constant temperature for several hours to optimize the reaction.

Many glasses in the systems \(M_2X\)–\(SiX_2\)–\(YX_n\)–\(MZ\), \(M = Li, Na, Ag; X = S, Se; Y = Ge, Al, B, P; Z = Cl, Br, I; n = 2, 3/2, 5/2\) were prepared in this manner [Kennedy and Zhang, 1988, Kennedy, 1989, Levasseur et al., 1981, Ribes et al., 1979].

However, thiosilicate glasses, even binary SiS₂ and SiSe₂ glasses, have a high tendency for crystallization. Therefore, faster quenching techniques were developed to produce glasses in larger (often, alkali-rich) composition domains. Several techniques can be considered from the simple crush of a droplet between two plates to more sophisticated techniques such as twin-roller quenching. In this technique, droplets of molten material are pushed between rotating twin rollers, giving 50–80 μm thick glassy flakes. The technique allows obtaining quenching rates of about \(10^6\) K/s [Pradel
et al., 1985]. Glasses of the system $x$Li$_2$S–$\left(1-x\right)$SiS$_2$ were prepared in this manner up to large modifier content ($x = 0.6$) [Pradel and Ribes, 1986]. Several series of thiosilicate and selenosilicate glasses were obtained in such a way [Aotani et al., 1994, Deshpande et al., 1988a,b, Michel-Lledos et al., 1992, Pradel et al., 1995, Pradel and Ribes, 1986, Tatsumisago et al., 1996].

An alternative way to prepare thiosilicate glasses, i.e. the mechanical milling technique, has been proposed by Morimoto in 1999 [Morimoto et al., 1999]. Powders of the starting (former, modifier …) compounds are placed in a jar with a series of balls. Mechanical milling treatment is carried out on the mixture using a planetary ball mill. The mechanical energy transferred to the powder due to collision with the balls allows the reaction between the components of the mixture. Optimization of different parameters such as number and size of the balls, rotation speed, allowed the synthesis of 0.6Li$_2$S–0.4SiS$_2$ fine powders with similar properties than the melt-quenched samples. An advantage of this method is its ability to proceed at room temperature.

Finally, thin films of several thiosilicate glasses, i.e. Li$_2$S–SiS$_2$, Li$_2$S–SiS$_2$–LiI, Li$_2$S–SiS$_2$–P$_2$S$_5$–LiI, were obtained by thermal evaporation of the bulk glasses, which required the development of a specially designed device installed in a glove box under a controlled argon atmosphere [Creus et al., 1989]. Amorphous films were obtained and further used as solid electrolytes in exploratory research for microbattery development [Creus et al., 1992]. Owing to the difficulties to prepare and handle thin films in controlled atmosphere, very few experiments of this type were carried out but one can note a recent work on thiogermainate glasses by Seo and co-workers who prepared films by magnetron sputtering of a glassy target in an Ar atmosphere [Seo et al., 2016].

3. **Network formers**

Any thiosilicate/selenosilicate glass comprises SiS$_2$/SiSe$_2$ as the network former compound.

In contrast to SiO$_2$ whose structure comprises a 3D network of corner-sharing (CS) tetrahedra (Td), whatever the considered polymorph (quartz, trydimite, cristobalite), SiS$_2$, in its unique known crystalline form, has a structure comprising infininte chains of edge-sharing (ES) tetrahedra (Figure 1) [Zintl and Loosen, 1935]. The situation is more complex in SiSe$_2$ since, in addition to the usually reported polymorph, (named 700-SiSe$_2$ hereafter) isotype of SiS$_2$ [Hillel and Cueilleron, 1971, Peters and Krebs, 1982, Weiss and Weiss, 1952], two other polymorphs, prepared by solid state reaction at 400 °C and 500 °C (named 400-SiS$_2$ and 500-SiSe$_2$ respectively), have been identified [Pradel et al., 1993].

The vitreous counterparts of crystalline SiS$_2$ and SiSe$_2$ (named SiS(e)$_2$ hereafter) can be prepared by conventional melt-quench technique even though they have a high tendency to crystallize. In the silicon sulfide and selenide families it is also possible to produce non-stoichiometric glasses, i.e. Si$_x$S$_{1-x}$ and Si$_x$Se$_{1-x}$, which is not the case in the oxide family counterpart [Tenhover et al., 1983a, Griffiths et al., 1984, Johnson, 1986]. This is based on the possibility
of forming homopolar S(e)–S(e) and Si–Si bonds. Indeed, in these families, the difference in electronegativity between Si (1.8) and S (2.6)/Se (2.6) is not so large, in contrast with the difference in electronegativity between Si (1.8) and O (3.5) in the oxide family. Therefore, homopolar bonds are not as disadvantaged as they are in oxide glasses.

It has to be noted that these differences in electronegativity has a strong consequence on the nature of the chemical bonds in the oxide and sulfide/selenide glasses. In oxide glasses, the chemical bonds have a strong ionic character, and SiO\textsubscript{2} can be considered as being built up with SiO\textsubscript{4}\textsuperscript{−} as the basic building block. In SiS(e)\textsubscript{2} glasses, the bonds have a strong ionocovalent character and the building blocks are usually rather written as SiS(e)\textsubscript{4/2}. Of course, both descriptions, i.e. SiO\textsubscript{4}\textsuperscript{−} and SiS(e)\textsubscript{4/2}, are just formal ones; they do not reflect the complex reality of the chemical bonds but clearly emphasize the difference between the two families.

Raman [Griffiths et al., 1984, 1985, Pradel et al., 1993, Sugai, 1987, Tenhover et al., 1983a,b, 1984, 1985] and \textsuperscript{29}Si NMR experiments [Moran et al., 1990, Pradel et al., 1993, Tenhover et al., 1988] as well as computer-modeling studies [Antonio et al., 1988, Gladden and Elliott, 1987, 1989a,b, Gladden, 1990] based on neutron-scattering data [Johnson et al., 1986a,b] were carried out in order to get an insight on the structure of the sulfide and selenide glasses and the unknown SiSe\textsubscript{2} polymorphs. \textsuperscript{29}Si NMR spectra of the SiSe\textsubscript{2} polymorphs and SiSe\textsubscript{2} glass are shown in Figure 2 [Pradel et al., 1993]. The NMR spectrum of the 700-SiSe\textsubscript{2} polymorph shows a single peak at −92.6 ppm which can be assigned to the unique Si site present in the structure and involved in ES tetrahedra. Similar findings were obtained for crystalline SiS\textsubscript{2} with a single \textsuperscript{29}Si NMR peak at −19.5 ppm [Tenhover et al., 1988]. The spectra of 400-SiSe\textsubscript{2} and 500-SiSe\textsubscript{2} polymorphs are more complex. The first one shows two peaks at −83.5 ppm and −63.0 ppm and the second one four peaks at −88.4 ppm, −61.6 ppm, −56.3 ppm and −27.3 ppm. The spectrum of glassy SiSe\textsubscript{2} shows three broad peaks at −86.0 ppm, −62.6 ppm, −28.7 ppm, similar to \textsuperscript{29}Si NMR spectrum of vitreous SiS\textsubscript{2} where the peaks appear at −16.8 ppm, −7.6 ppm and +7.5 ppm [Tenhover et al., 1988]. The presence of several peaks in the \textsuperscript{29}Si NMR spectra of SiS(e)\textsubscript{2} compounds suggests the existence of several Si environments. Three possible Si environments have been suggested: Si belonging to tetrahedra sharing two, one or zero edge(s), shown in Figure 3 and labeled E2, E1 and E0 respectively [Antonio et al., 1988, Gladden and Elliott, 1987, 1989a,b, Gladden, 1990, Griffiths et al., 1984, 1985, Moran et al., 1990, Pradel et al., 1993, Sugai, 1987, Tenhover et al., 1983a,b, 1984, 1985, 1988]. For selenide (sulfide) compounds, peaks in the region −80/−90 ppm (−10/20 ppm) are attributed to E2 species, those in the region −60/−70 ppm (−7/8 ppm) to E1 species and E0 species have their signature in the region −20/−30 ppm (+7/+8 ppm). Therefore, 400-SiSe\textsubscript{2} polymorph comprises E2 and E1 species in the ratio 1/1. A tentative structural model that fulfills this requirement is proposed in Figure 1 [Pradel et al., 1993]. Both, 500-SiSe\textsubscript{2} and vitreous SiSe\textsubscript{2} comprise the three species E2, E1 and E0, suggesting a close structural relationship between them.
However, a minor discrepancy between E2:E1:E0 ratios (1:2:2 for the crystalline compound compared to 1:2:1 for the glass) exists, so it was suggested that an even better agreement could be achieved if one considered that the glass was comprised of a mixture of the units existing in 400-SiSe$_2$ and 500-SiSe$_2$. It was supported by the fact that the vitreous transition temperature, i.e. the temperature at which the vitreous structure is frozen in, lies around 450 °C. All Raman and computer-modeling studies based on neutron diffraction experiments (not described in detail here) are consistent with the above findings, i.e. they confirm the coexistence of E2, E1 and E0 species [Antonio et al., 1988, Gladden and Elliott, 1987, 1989a,b, Gladden, 1990, Griffiths et al., 1984, 1985, Moran et al., 1990, Pradel et al., 1993, Sugai, 1987, Tenhover et al., 1983a,b, 1984, 1985, 1988]. On the other hand, they all failed in one way or another to give a complete description of the glass structure that agrees with all experimental data (relative amounts of species for example). Some models such as the random network model by Sugai [Sugai, 1986, 1987], are too simplistic. In the cross-linked-chain-cluster (CLCC) model, Griffiths et al. proposed an intermediate range order built up with chain fragments containing E2 units interconnected by E1 and E0 tetrahedra, as shown in Figure 4 [Griffiths et al., 1984]. This model was refined by Gladden in an investigation based upon a computer-modeling study of neutron diffraction data [Gladden and Elliott, 1987, 1989a]. In this work, several types of intermediate range order species were considered, including random chains and the Griffiths CLCC species. The best agreement led to a mixture of 85% random chains containing an average of seven E2 units and 15% of CCLC units, consisting of rings with two E0 and four E2 units.

Figure 3. Possible Si environments in alkali silicate, thiosilicate and selenosilicate compounds.
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Figure 4. Schematic representation of a cross-linked chain cluster (CLCC) proposed by Griffiths and collaborators [Griffiths et al., 1984].

4. Network former/network modifier

4.1. Single modifier

In oxide glasses, addition of a modifier oxide, e.g. \( \text{Li}_2\text{O}, \text{Na}_2\text{O} \), in the former oxide \( \text{SiO}_2 \), creates the well-known non-bridging oxygen \( \text{(NBO)} \) and the consequent \( \text{Q}_n \) species of tetrahedrally coordinated silicon bound to a number, \( n \), of bridging oxygen \( \text{(BO)} \) \( (n = 4, 3, 2, 1 \text{ or } 0) \) as shown in Figure 3. Do similar non-bridging sulfur or selenium \( \text{(NBS)} \) exist when a modifier is added to \( \text{SiS}_2 \)? In such a case, owing to the presence of ES tetrahedra, a larger number of entities, summarized in Figure 3, might be expected. For example, a \( \text{E1Q3} \) entity would be a tetrahedron with three bridging sulfur or selenium \( \text{(BS)} \), one NBS and sharing an edge with its neighbor.

Structural investigation of glasses \( \text{M}_2\text{X–SiX}_2 \) \( (\text{M} = \text{Li, Na, Ag to a less extent; X = S, Se}) \) was carried out using many complementary characterization techniques including Raman, IR, \( ^{29}\text{Si} \), \( ^{23}\text{Na} \), \( ^{7}\text{Li} \) NMR [Pradel et al., 1995], XPS [Foix et al., 2001, 2006] and neutron diffraction [Lee et al., 1997]. No technique alone could give a complete pattern of the structure at local and intermediate orders. However, their combination helped in getting a clear, even though still incomplete picture.

Information on glass structure can be obtained by comparing data obtained for glasses with those for crystalline model compounds. On this basis, crystallized thiosilicate compounds in the system \( \text{Na}_2\text{S–SiS}_2 \), where crystallographic data are numerous [Cade et al., 1972, Olivier-Fourcade et al., 1972, 1978], have been used to understand \( ^{29}\text{Si} \) NMR spectra of \( \text{Na}_2\text{S–SiS}_2, \text{Li}_2\text{S–SiS}_2 \) and \( \text{Ag}_2\text{S–SiS}_2 \) glasses [Eckert et al., 1989, Pradel et al., 1995]. The main information of this investigation can be summarized as follows. For all of the alkali-containing system, the spectra are dominated by two principal resonances in the chemical shift regions 1–7 ppm and
−7 to −12 ppm, as shown in Figure 5 for the Li$_2$S–SiS$_2$ system. The first one is the signature of E0 entities, with no edge shared with neighbors; the second is the signature of E1 entities with a single edge shared with neighbor. At low modifier content (10% Li$_2$S), still remains some E2 entities, whose signature occurs at −20 ppm. Addition of modifier results in the preferential destruction of ES tetrahedra, this tendency being much stronger for Li$_2$S than for Na$_2$S, 23% E1 remaining in 0.5Li$_2$S–0.5SiS$_2$ against 50% E1 (48% according to Watson and Martin [2017]) in 0.5Na$_2$S–0.5SiS$_2$. Note that no ES tetrahedra exists in the related Ag$_2$S–SiS$_2$ glasses [Pradel et al., 1995].

Owing to the very small $^{29}$Si chemical shift variations for thiosilicates, $^{29}$Si NMR cannot give direct and straightforward information on BS and NBS presence and distribution. In a recent re-investigation of the structure of Na thiosilicate glasses, complementary deconvolution of both $^{29}$Si NMR and Raman spectra was used to extract this information [Watson and Martin, 2017]. XPS experiments have also been carried out to study Li and Na thiosilicate glasses [Foix et al., 2001]. Data from XPS core peak analysis for lithium glasses are shown in Figure 6. For all glasses (the same applies for Na glasses), a decomposition of the large S$_{2p}$ peak into a minimum of two doublets was necessary in order to fit the experimental curve. These doublets are the signature of BS and NBS, similar to the BO and NBO in oxide glasses. The signal at lower energy was assigned to NBS that are more negative than BS associated with the doublet at higher energy. When the modifier increases, the proportion of NBS simultaneously increases in the glass.

**Figure 5.** $^{29}$Si MAS NMR spectra for $x$Li$_2$S–$(1-x)$SiS$_2$ glasses.

**Figure 6.** XPS S$_{2p}$ core peaks for $x$Li$_2$S–$(1-x)$SiS$_2$ glasses. Doublet 1: NBS; doublet 2: BS.
A thorough analysis of the XPS data including comparison with XPS spectra of model crystalline compounds has been carried out. Based on these data, Li$_2$SiS$_3$ and Na$_2$SiS$_3$ clusters were built and a Mulliken population analysis was carried out to investigate the charge distribution of the clusters. The main result of the whole study is the noticeable difference existing between the structures of lithium and sodium thiosilicate glasses, in agreement with NMR results. According to XPS data, this is due to different electronic redistribution over the network when one or the other alkali is added, the sodium addition resulting in a change in the electronic distribution over the entire network, which also affects BS. In oxide glasses, the splitting of 2 eV between NBO and BO is larger than the splitting of $\sim$0.9 eV between NBS and BS in sulfide glasses. It can be explained by the decreased ionic nature of the bonds in thiosilicate and smaller difference in the electronic density around NB and B atoms. XPS valence band spectra of Li$_2$SiS$_3$ and Na$_2$SiS$_3$ glasses combined with theoretical calculations have also been carried out [Foix et al., 2006]. The difference in spectra of Li and Na thiosilicate glasses points for the presence of a majority of CS tetrahedra in the Li glasses whereas a larger proportion of ES tetrahedra should be present in the Na glass, in agreement with $^{29}$Si NMR studies.

Structural investigation of lithium and sodium selenosilicate glasses and crystalline model compounds was also carried out using mainly Raman and $^{29}$Si NMR experiments [Pradel et al., 1992, 1995]. $^{29}$Si NMR proved to be very powerful in this case. Indeed, much larger $^{29}$Si chemical shifts were observed as compared to sulfide glasses and, therefore, species with different Qn could be discriminated as shown in Figure 7 for the Li$_2$Se–SiSe$_2$ family. It can be seen that the tendency to maintain ES species is lower in selenosilicate glasses than in thiosilicate ones. Indeed, none are left at $x=0.5$ for selenosilicates. The combination of absence of ES species and possible discrimination between Qn species gave the opportunity to test different structural models that were much discussed in oxide glasses to identify the population distribution of Qn species, e.g. a completely random “statistical” [Schramm et al., 1984] or chemically ordered [Dupree et al., 1984, Grimmer et al., 1984] distribution. Figure 8 shows the experimental spectrum of 0.6Na$_2$Se–0.4SiSe$_2$ glass and the resulting simulated distribution based upon the experimental chemical shifts of Qn sites. The two extreme scenarios can be discarded: While the binary model predicts the presence of Q1 species only, the random model would predict less Q1 species than observed.

On the whole, the structural investigation showed a more complex structure in thio and selenosilicate glasses than in oxide analogs. On the other hand, as it occurs in oxide glasses, the addition of modifier leads to the creation of non-bridging atoms (here S or Se ones), even though the Si–S(e) bonds are less ionic than the Si–O ones. The nature of both the modifier and chalcogen have consequences on different aspects of the structure such as the number of ES species and the charge on the atoms.

Figure 7. $^{29}$Si MAS NMR spectra for $x$Li$_2$Se–(1–$x$)SiSe$_2$ glasses. Spinning sidebands are indicated by asterisks.
Physicochemical characterization of these glasses has been carried out in order to understand the influence of the structure on glass properties. Figure 9 shows the evolution of the glass transition temperature $T_g$ with the addition of modifier in glasses $xM_2X-(1-x)SiX_2$ with $M = Li, Na, Ag$ and $X = S, Se$ [Pradel and Ribes, 1989a, Michel-Lledos et al., 1992, Tenhover et al., 1983a,b, 1984, Johnson, 1986]. $T_g$ decreases rapidly with the first addition of modifier, then at a slower pace, if any, to further addition. A rather large difference between $T_g$ of the sulfide and selenide systems could partly be explained by their structural difference i.e. the larger presence of ES species in the sulfide glasses [Pradel and Ribes, 1992]. The electrical characteristics (electrical conductivity and activation energy) of $xM_2X-(1-x)SiX_2$ glasses with $M = Li, Na, Ag$ and $X = S, Se$ have been investigated. As an example, Figure 10 shows the composition dependence of room temperature conductivity and activation energy for lithium-conducting glasses including oxide ones [Pradel and Ribes, 1992, Yoshiyagawa and Tomozawa, 1982]. Sulfide and selenide glasses have comparable electrical characteristics. In contrast, the corresponding oxide glasses have much lower conductivities and much larger activation energies. This can be understood if the main factor that controls ion motion is the anion polarizability, which is comparable for S (7.3) and Se (7.5) and much lower for O (3.1). As described in the introduction section, this was predicted by the weak electrolyte theory, which is at the basis of the investigation of these glasses. At this point, another family of thiosilicate glasses, which will not be discussed further in the paper, can be mentioned, i.e. glasses with a “salt”, e.g. a lithium halide, dissolved in the matrix [Akridge, 1984, Sahami et al., 1985, Kennedy et al., 1986, Pradel and Ribes, 1989a]. These glasses show increased conductivity owing to the high polarizability of the anions and increase in numbers of mobile ions. Up to 40% of salt can be dissolved in a glass but it is at the expense of its thermal stability.

4.2. Multi-modifiers

In the previous section, it has been shown that addition of Li$_2$S or Na$_2$S to SiS$_2$ has somewhat different consequences on the glass structure and properties with a stronger tendency for Li$^+$ ions to destroy the ES tetrahedra and a charge redistribution over the entire network for Na$^+$ ions leading to more negative BS and NBS. The question is: what would happen if both Li$^+$ and Na$^+$ were introduced in the glass network? Some answers were obtained with the investigation of the series of glasses 0.5$(xNa_2S-(1-x)Li_2S)$–0.5SiS$_2$
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Figure 10. Conductivity $\sigma$ and activation energy of conductivity $E_\sigma$ for $x\text{Li}_2\text{X}-(1-x)\text{SiX}_2$ glasses ($\text{X} = \text{O}, \text{S} \text{and Se}$).

with $0 \leq x \leq 1$. The structural investigation was only carried out by $^{29}\text{Si}$ NMR [Pradel et al., 1995]. It was shown that, while the two main peaks observed in the binary glasses and related to $E_0$ and $E_1$ species (Figure 5) were still present in all mixed Na/Li glasses, their relative intensity was closer to that observed for the binary lithium glasses, even for the first addition of lithium in the sodium glass (Figure 11). The chemical shifts for the two peaks were intermediate between those measured for the binary glasses, which led to the conclusion of the absence of any preferential association of one type of cations for a specific site $E_1$ or $E_2$.

While the replacement of lithium by sodium in the thiosilicate glasses does not seem to transform the structure drastically, it has drastic consequences on several properties. Dependence of glass transition temperature, $T_g$, electrical characteristics (conductivity $\sigma$ and activation energy of conductivity $E_\sigma$) and activation energy of the relaxation time $T_1$, $E_{T_1}$, (measured by $^7\text{Li}$ NMR spectroscopy) on composition are shown in Figure 12 [Pradel and Ribes, 1994]. A strong non-linear variation of these characteristics is observed. $T_g$ and $\sigma$ show a high minimum at $x \sim 0.5$, whereas the activation energies $E_\sigma$ and $E_{T_1}$ display a maximum for the same values of $x$. These features are the signature of the mixed alkali effect (MAE), a very well-known phenomenon, reported many times in oxide glasses [Day, 1976, Ingram, 1987, 1992], in particular in silicate glasses, the oxide counterparts of the glasses described here [Elliott, 1992]. MAE occurs when a modifier is substituted by another one, at a constant total modifier content. Therefore, it appears to be a common feature between oxide and chalcogenide glasses, even though reports on MAE in the second family are scarce and concern mainly two systems, the present one and the $\text{Ag}_2\text{S}$–$\text{Rb}_2\text{S}$–$\text{GeS}_2$ system (here, we should rather call it “mixed cation effect”) [Rau et al., 2001].

The information reported above in Section 4 indicates that $\text{Li}^+$ and $\text{Na}^+$ ions do not favor the same sites in thiosilicate glasses; in particular, NBS are more negative around $\text{Na}^+$ than in the neighborhood.
of Li$^+$. In the Ag$_2$S–Rb$_2$S–GeS$_2$ system, it has been clearly demonstrated by EXAFS experiments that the Ag$^+$ and Rb$^+$ cations maintained their own environment in the mixed glasses. A similar situation probably applies to the Li/Na thiosilicate glasses. So it can be anticipated that for an ion, to hop from one site to the next is more difficult in a mixed environment than in single alkali one where all the sites are similar and match the only mobile ion. The mobility would then be restricted, which would induce a lower conductivity and higher activation energy as observed.

Figure 12. (a) Conductivity $\sigma$ and activation energy of conductivity $E_\sigma$ and (b) glass transition temperature, $T_g$ and activation energy of the relaxation time $T_1$, $E_{T_1}$ for 0.5$[x$Na$_2$S–(1 – x)Li$_2$S]–0.5SiS$_2$ glasses. The dashed lines are guidelines for experimental data.

Figure 13. Glass formation region in the Li$_2$S–Al$_2$S$_3$–SiS$_2$ system. Circles and triangles correspond to glasses, while stars refer to partially crystallized samples.

5. Effect of a competitive network former on thiosilicate glasses

5.1. Thio-aluminosilicate

The glass forming ability of compositions belonging to the Li$_2$S–Al$_2$S$_3$–SiS$_2$ system, sulfide counterparts of lithium aluminosilicates, has been investigated [Deshpande et al., 1988b, Hayashi et al., 2004, Martin and Sills, 1991, Pradel and Ribes, 1992]. The explored region and vitreous domain extension, which depends upon the elaboration technique, i.e. conventional melt-quenching, twin-roller quenching and mechanical alloying, is shown in Figure 13. Compared to the oxide family, the glass forming ability is poor with a maximum ratio of Al/Li much below the 1.5 reported for oxide glasses.

Raman and $^{29}$Si, $^{27}$Al NMR investigations helped in getting some insights into the structure of these glasses. Raman spectra of glasses 0.5Li$_2$S–xAl$_2$S$_3$–(0.5 – x)SiS$_2$ with $x = 0, 0.1, 0.25$, along with the spectra of crystalline SiS$_2$ and crystalline $\alpha$–Al$_2$S$_3$ are shown in Figure 14 [Pradel and Ribes, 1992]. The low temperature form of crystalline Al$_2$S$_3$, $\alpha$–Al$_2$S$_3$, has a lacunar wurtzite structure with aluminum occupying tetrahedral sites, a difference with the high temperature form of crystalline Al$_2$O$_3$ where aluminum is in an octahedral environment. The main band in the Raman spectrum at 248 cm$^{-1}$ is assigned to the symmetric valence vibrations of the AlS$_{4/2}$
Figure 14. Raman spectra of 0.5 Li$_2$S–xAl$_2$S$_3$–(0.5 – x)SiS$_2$ glasses, (Si–S) and (Si–S$^-$) stand for bridging Si–S bonds and non-bridging Si–S$^-$ bonds, respectively. Raman spectra of crystalline SiS$_2$ and α–Al$_2$S$_3$ are reported for comparison.

Figure 15. Conductivity $\sigma$, activation energy of conductivity $E_\sigma$ and glass transition temperature $T_g$ of 0.5Li$_2$S–xAl$_2$S$_3$–(0.5 – x)SiS$_2$ glasses elaborated by twin-roller quenching. The dashed lines are guidelines for experimental data.

attributed to the stretching vibrations of Al–S bonds. Compared to the crystalline compound, the band attributed to Al–S bonds is displaced toward high frequencies, consistent with a shortening of the bonds and therefore, leading to discarding an octahedral coordination for the aluminum. It is in agreement with the results from $^{27}$Al NMR experiments [Martin and Sills, 1991]. In this case, the NMR spectra of all prepared Li$_2$S–Al$_2$S$_3$–SiS$_2$ glasses show a single peak at 110 ppm, similar to the peak present in the NMR spectrum of crystalline α-Al$_2$S$_3$. In this case, one expects the lithium to act as a charge compensator and therefore, a decrease in the number of non-bridging sulfur, consistent with the decrease in the band at 390 cm$^{-1}$ in the Raman spectra.

This assumption is supported by the increase of the glass transition temperature $T_g$ with the Al$_2$S$_3$ content in the 0.5Li$_2$S–xAl$_2$S$_3$–(0.5 – x)SiS$_2$ glasses as shown in Figure 15. It is consistent with an increasing polymerization of the network due to the disappearance of non-bridging sulfur. Such an increase of $T_g$ with increase in the Al$_2$S$_3$ content in the glasses has been reported for other series of compositions such as series where the ratio Li/Al was constant and equals to 2/1, 1/1, 1/2 [Martin and Sills, 1991].

The electrical properties of Li$_2$S–Al$_2$S$_3$–SiS$_2$ glasses have been reported. The conductivity $\sigma$ and activation energy of conductivity $E_\sigma$ for the
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5.2. Thiogermanosilicate

The structure of Si₄Ge₁₋ₓS₂ glasses has been investigated by analyzing Raman experiments [Tenhover et al., 1983b]. It was shown that, whatever be x, separate clusters of the two formers were obtained, which was attributed to the impossible mixing of SiS₂ and GeS₂ networks that adopt different medium-range order configurations: dominance of ES tetrahedra as shown in Section 3 for the first one and dominance of CS tetrahedra for the second one [Feltz et al., 1985].

The glass forming ability and properties of glasses, whose composition lies along two lines of the system Li₂S–SiS₂–GeS₂, i.e. 0.3Li₂S–0.7[(1 – x)SiS₂–xGeS₂] and 0.5Li₂S–0.5[(1 – x)SiS₂–xGeS₂] with 0 ≤ x ≤ 1, were also investigated (Figure 16) [Deshpande et al., 1988a, Pradel and Ribes, 1989b, Pradel et al., 1998]. The glass elaboration required the use of the twin-roller quenching technique. Main structural information was obtained by a Raman experiment analysis. Raman spectra of 0.3Li₂S–0.7[(1 – x)SiS₂–xGeS₂] glasses. (Si(Ge)–S) and (Si(Ge)–S⁻) stand for bridging Si(Ge)–S bonds and non-bridging Si(Ge)–S⁻ bonds, respectively. Blue dashed line corresponds to the main peak of 0.5Li₂S–0.5SiS₂ glass. Raman spectrum of crystalline GeS₂ is reported for comparison. SAXS intensity for five glasses belonging to the three different regions is shown in the inset.

Figure 16. Investigated glass compositions in the system Li₂S–SiS₂–GeS₂, i.e. 0.3Li₂S–0.7[(1 – x)SiS₂–xGeS₂] and 0.5Li₂S–0.5[(1 – x)SiS₂–xGeS₂].

0.5Li₂S–xAl₂S₃–(0.5 – x)SiS₂ glasses are shown in Figure 15. The slight decrease in σ and increase in Eσ with the addition of Al₂S₃ point for a lower mobility of the lithium acting as charge compensator as compared to lithium close to a non-bridging sulfur. Such assumption is supported by the fact that, in the series yLi₂S–0.1Al₂S₃–(0.9 – y)SiS₂, the conductivity increases from 3.3 × 10⁻⁶ S·cm⁻¹ to 1.3 × 10⁻⁴ S·cm⁻¹ when y changes from 0.3 to 0.6.

Figure 17. Raman spectra of 0.3Li₂S–0.7[(1 – x)SiS₂–xGeS₂] glasses. (Si(Ge)–S) and (Si(Ge)–S⁻) stand for bridging Si(Ge)–S bonds and non-bridging Si(Ge)–S⁻ bonds, respectively. Blue dashed line corresponds to the main peak of 0.5Li₂S–0.5SiS₂ glass. Raman spectrum of crystalline GeS₂ is reported for comparison. SAXS intensity for five glasses belonging to the three different regions is shown in the inset.

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Figure 18. Glass transition temperature, $T_g$, density, $\rho$, for Li$_2$S–SiS$_2$–GeS$_2$ glasses. Filled and empty symbols correspond to series 0.3Li$_2$S–0.7[(1–x)SiS$_2$–xGeS$_2$] and 0.5Li$_2$S–0.5[(1–x)SiS$_2$–xGeS$_2$], respectively. The inset displays the DSC curve of 0.3Li$_2$S–0.25SiS$_2$–0.45GeS$_2$ glass. The dashed lines are guidelines for experimental data.

Figure 19. Conductivity $\sigma$ and activation energy of conductivity $E_\sigma$ for Li$_2$S–SiS$_2$–GeS$_2$ glasses. Filled and empty symbols correspond to series 0.3Li$_2$S–0.7[(1–x)SiS$_2$–xGeS$_2$] and 0.5Li$_2$S–0.5[(1–x)SiS$_2$–xGeS$_2$], respectively. The dashed lines are guidelines for experimental data.

non-bridging Si–S$^-$ (Ge–S$^-$) bonds at 412 cm$^{-1}$ (338 cm$^{-1}$) and 368 cm$^{-1}$ (395 cm$^{-1}$) respectively [Souquet et al., 1981]. When considering the general evolution of the spectra, three different domains can be identified: For the first substitutions of a glass former by the other one, i.e. for $0 \leq x < 0.50$ and $0.64 < x \leq 1$, a smooth change in the spectra is observed. A sudden change occurs when the Si/Ge ratio is close to 1, i.e. for $0.5 \leq x \leq 0.64$. For $x = 0.5$ the spectrum narrows and the main peak suddenly shifts to higher frequency, close to the frequency of the stretching vibrations of non-bridging Si–S$^-$ bonds in 0.5Li$_2$S–0.5SiS$_2$ glass (377 cm$^{-1}$). The two spectra look very much alike, but with a broadening for the mixed glass due to the presence of Ge–S bonds, which also explains the further broadening of the spectra upon further GeS$_2$ addition. The situation is different in the second series of mixed glasses, i.e. 0.5Li$_2$S–0.5[(1–x)SiS$_2$–xGeS$_2$]. In this case, a smooth evolution of the Raman spectra (not shown here, Deshpande et al. [1988a]) is observed all along the line from $x = 0$ to $x = 1$. SAXS experiment on the first series 0.3Li$_2$S–0.7[(1–x)SiS$_2$–xGeS$_2$] were carried out in order to check the homogeneity of the glasses [Pradel et al., 1998]. Samples from the three regions were tested. Typical results are shown in the inset of Figure 17. Data obtained for samples belonging to the limiting regions ($0 \leq x < 0.50$ and $0.64 < x \leq 1$) could be fitted in the framework of the Debye–Buëche model [Debye and Buëche, 1949] and therefore, are homogeneous. In contrast, the higher scattering observed for glasses belonging to the central region ($0.5 \leq x \leq 0.64$) rather obeys Porod’s law [Porod, 1982], which indicates the presence of aggregates (of about 50 Å) in the glass. Both Raman and SAXS experiments point toward homogeneous glasses in the limiting regions and the occurrence of a phase separation in the central one, leading probably to entities with a composition close to GeS$_2$ embedded in a matrix whose composition is close to that of 0.5Li$_2$S–0.5SiS$_2$ glass, i.e. Li$_2$SiS$_3$. 

Vitreous transition temperatures $T_g$ and density $\rho$ were measured and are shown in Figure 18 while the electrical conductivity $\sigma$ and corresponding activation energy $E_\sigma$ are shown in Figure 19. A smooth evolution of $T_g$, $\sigma$ and $E_\sigma$ when SiS$_2$ is substituted by GeS$_2$ is observed for the 0.5Li$_2$S–0.5[(1–x)SiS$_2$–xGeS$_2$] glasses, which agrees with Raman data. In contrast, sudden changes in all parameters $\rho$, $T_g$, $\sigma$ and $E_\sigma$ occur in the central region for compositions pointed as phase-separated from SAXS and Raman
experiments. Appearance of two $T_g$ and lowering of $\rho$ strongly support the presence of a phase separation. Moreover, the larger conductivity of the glasses in the central region, close to the conductivity of the 0.5Li$_2$S–0.5SiS$_2$ glass, supports the structural scheme suggested by Raman data, i.e. GeS$_2$ aggregates embedded in a Li$_2$SiS$_3$ matrix, provided that a percolation threshold is reached.

The differences between the two families of glasses, homogeneity versus phase separation, could be understood by the degree of modification of the matrix, which, upon increase, tends to lead to similar configurations for the two end-line compounds. GeS$_2$ favors CS tetrahedra, hence few, if any, ES tetrahedra are expected in lithium thiogerminate glasses. For 0.3Li$_2$S–0.7SiS$_2$ glass, 34% of tetrahedra in the structure are ES ones; this number goes down to 23% for 0.5Li$_2$S–0.5SiS$_2$ glass [Eckert et al., 1989]. Therefore, mixing of the two end-line compounds is more favorable at high modifier content. It can also be pointed out that a crystalline metathiosilicate phase, Li$_2$SiS$_3$, exists [Weiss and Rocktäschel, 1960], whereas crystalline thiodisilicate, Li$_2$Si$_2$S$_5$ (0.33Li$_2$S–0.67SiS$_2$), does not. The phase separation occurs in the 0.3Li$_2$S–0.7[(1–x)SiS$_2$–xGeS$_2$] with the tendency to produce a glassy phase of composition Li$_2$SiS$_3$. On the other hand, homogeneous glasses are obtained for the family 0.5Li$_2$S–0.5[(1–x)SiS$_2$–xGeS$_2$] corresponding to the Li$_2$SiS$_3$ composition. This can be related to an investigation on lithium germanosilicate glasses, the oxide counterparts of the present glasses [Verweij et al., 1979]. Anomaly in the refractive index of several Li$_2$O–SiO$_2$–GeO$_2$ glasses was indeed explained by the tendency of glasses to phase-separate with one of the resulting phase having the disilicate composition. In contrast, no anomaly was observed for 0.33Li$_2$O–0.67[(1–x)SiO$_2$–xGeO$_2$] glasses corresponding to the exact disilicate composition. A similar situation to ours since, in the oxide glasses, crystalline disilicate phase exists while the metasilicate one does not.

5.3. Thiophosphosilicate

Investigation of lithium thiophosphosilicate glasses of composition 0.6Li$_2$S–0.4[(1–x)SiS$_2$–xP$_2$S$_5$] with 0 ≤ x ≤ 1 was carried out in the eighties (Figure 20) [Kennedy and Zhang, 1989, Kennedy et al., 1990]. The main information is the anomalous evolution of some properties, e.g. glass transition temperature, when substituting a modifier by the other one, with a change at the particular composition corresponding to a ratio SiS$_2$:P$_2$S$_5$ equals to 2:1, i.e. Si/P = 1. Even though a paper, mainly focussed on electrochemical properties of this lithium family, was published very recently [Zhao et al., 2020], the renewal of interest for the thiophosphosilicate glasses concerns mainly the sodium family, and matches the increasing interest for the development of Na all-solid-state batteries. Two series of glasses with composition 0.5Na$_2$S–0.5[xSiS$_2$–(1–x)P$_5$S$_5$/2] and 0.67Na$_2$S–0.33[xSiS$_2$–(1–x)P$_5$S$_5$/2] (purple symbols) and line of investigated compositions in the Li$_2$S–SiS$_2$–P$_2$S$_5$ system, i.e. glasses with 60 mol% Li$_2$S.

![Figure 20. Investigated glass compositions in the system Na$_2$S–SiS$_2$–PS$_5$/2, i.e. 0.5Na$_2$S–0.5[xSiS$_2$–(1–x)P$_5$S$_5$/2] and 0.67Na$_2$S–0.33[xSiS$_2$–(1–x)P$_5$S$_5$/2] (purple symbols) and line of investigated compositions in the Li$_2$S–SiS$_2$–P$_2$S$_5$ system, i.e. glasses with 60 mol% Li$_2$S.](image-url)
presence of both P Q0 and Si Q3 in glasses comprising 50 mol% Na2S indicates that there is unequal sharing of the Na⁺ ions between the two glass formers; P taking additional Na⁺ to form Q0 units and Si giving up Na⁺ to form Q3 structures. For the 0.67Na2S–0.33[xSiS2–(1 – x)PS5/2] glasses, some complexity is added to the structure since a significant concentration of unreacted Na2S species are present in all glasses. Densities ρ and glass transition temperatures \( T_g \) were measured for all glasses (Watson, 2018b). Their evolution with composition is consistent with the structural models proposed for the glasses. As an example, Figure 22 shows the evolution of \( T_g \), molar volume (calculated from experimental ρ) and fraction of BS (BS/(BS + NBS) calculated from the distribution of all species) when P is substituted by Si for 0.5Na2S–0.5[xSiS2–(1 – x)PS5/2] glasses. \( T_g \) remains fairly constant for the first additions of SiS2 when the structure is still dominated by P Q1 and Q0 species (Figure 21). When \( x \geq 0.5 \), Si Q2 and Q3 species starts being dominant and hence, the fraction of BS. Then, \( T_g \) starts and keeps on increasing with further addition of SiS2. At the opposite, the molar volumes decrease with the increase in BS and therefore, with the networking increase through the increase in Si Q2 and Q3 species. Finally, a 23Na NMR investigation of these glasses has been reported recently [Shastri et al., 2019], which gives information on the dynamics and distribution of Na⁺ ions. The most relevant information, suggested by a Na NMR second moments study and related to the glass structure, is a homogeneous distribution of sodium in the glass network over most of the composition range, but for Na-richest ones where sodium tend to cluster.

5.4. Thioborosilicate

As far as we know, a single composition, i.e. the 0.30Li2S–0.21SiS2–0.09B2S3–0.40LiI glass, has been investigated in the late eighties [Kennedy, 1989]. Nothing had been done in these systems ever since, until very recently a structural investigation of a series of glasses with composition 0.6Na2S–0.4[(1 – x)SiS2–xBSiS3/2], 0 ≤ x ≤ 1, has been reported (inset Figure 23) [Curtis et al., 2019]. IR, Raman, 29Si and 11B NMR experiments were carried out in order to study the changes occurring in the glass structure when a glass former was substituted by the other one.

The structural analysis was similar to the analysis described in the previous section for thiophosphosilicate glasses. In the same way, deconvolution of Raman and NMR spectra helped in proposing a picture of the glass structure on the basis of the distribution of the different species likely to exist in the glass, and resulting from disproportionation reaction.
Figure 23. Composition dependence of the fractions of the main $\text{B Q}_n$ and $\text{Si Q}_n$ units in $0.6\text{Na}_2\text{S} + 0.4[\text{xBS}_{3/2} + (1-x)\text{SiS}_2]$ glasses. In the inset is shown the line of studied compositions.

The different species for these thioborosilicate glasses and their distribution are shown in Figure 23. In fact, owing to the difficulty to obtain pure $\text{B}_2\text{S}_3$ exempt of oxygen, several oxi-sulfide species were present in the glasses. These minority species are not shown in Figure 23. The main information can be summarized as follows. Tetrahedrally coordinated $\text{Si}$ with a single $\text{BS}$ (Si Q1) are the dominant species in the end-line member, $0.6\text{Na}_2\text{S}–0.4\text{BS}_{3/2}$ glass while $\text{BQ}_0$ corresponding to 3-coordinated boron with no BS is the dominant one in $0.6\text{Na}_2\text{S}–0.4\text{BS}_{3/2}$ glass. Upon substitution, there is a disproportionate sharing of the $\text{Na}^+$ between the $\text{B}$ and $\text{Si}$ subnetworks with $\text{Si}$ taking additional $\text{Na}^+$ to form $\text{Si Q}_0$ entities and $\text{B}$ giving up $\text{Na}^+$ to form $\text{B Q}_1$ species. Once again, it is interesting to note that no crystalline phase with composition $\text{Na}_6\text{Si}_2\text{S}_7$ has ever been reported, while $\text{Na}_4\text{SiS}_4$ exists [Cade et al., 1972, Eckert et al., 1989]. On the whole, it appears that, in multi-former thiosilicate glasses, the thiosilicate subnetwork tends to adopt a configuration close to that of a neighboring crystalline phase.

6. Conclusion

Thio- and selenosilicate glasses are the sulfide and selenide analogs of silicate glasses. Owing to the different characteristics of the chalcogen, e.g. size and polarizability, these glasses show markedly different structures and properties, in particular, a much higher electrical conductivity $\sigma$, which attracted attention for the development of lithium and more recently sodium all-solid-state batteries. The structure of the glass formers, $\text{Si(S(e))}_2$, is comprised of both ES and CS tetrahedra, a major difference with silica. Upon addition of a modifier, a depolymerization of the network with appearance of NBS occurs, which is similar to the creation of NBO in silicate glasses. Preferential destruction of ES tetrahedra is generally observed with an increasing trend in the sequences: S to Se and Na to Li to Ag. Thiosilicate glasses comprising two modifiers show the very well-known mixed alkali effect with a strong decrease in both, $T_g$ and $\sigma$ for mixed glasses compared to single ones, a common characteristic with silicates. When added to lithium thiosilicate, aluminum sulfide, $\text{Al}_2\text{S}_3$, behaves as a former compound, which decreases the number of NBS and results in an increase in $T_g$ and slight decrease in $\sigma$. The mixed glass former effect, observed when a glass former is replaced by another one, the total amount of modifier being constant, has been investigated in thiosilicate glasses with the second former compound being $\text{GeS}_2$, $\text{PS}_{5/2}$ or $\text{BS}_{3/2}$. Depending upon the amount of modifier, lithium thiogermanosilicate glasses are either homogeneous or phase-separated, with a nonlinear variation of properties, e.g. $\rho$, $T_g$ and $\sigma$, in the second case. A trend to phase-separate and produce a subnetwork with composition of an existing crystalline phase has also been reported in the oxide analogs, the germanosilicate glasses. In sodium thiophosphosilicate and thioborosilicate glasses, disproportionation reactions between the two former occur with the transfer of $\text{Na}^+$ from the Si subnetwork to the P subnetwork in the first family. In contrast, $\text{Na}^+$ are transferred from the B subnetwork toward the Si subnetwork in thioborosilicate glasses. In both case, the resulting composition of the Si subnetwork tends to acquire the composition of an existing crystalline phase, i.e. $\text{Na}_4\text{Si}_4\text{S}_{10}$ corresponding to $\text{Si Q}_3$ species and $\text{Na}_4\text{SiS}_4$ corresponding to $\text{Si Q}_0$ species.
One of the main interests of the investigation of these thio- and selenosilicate glasses is to shed light on the consequence of a change in the nature of the chemical bonds, from highly ionic in oxides to strongly ionocovalent in sulfides/selenides, on the structure and physicochemical properties of the obtained glasses. Apart from this academic interest, these glasses could get renewal of interest since they could be at the basis of the elaboration of new glass-ceramics to be used as solid electrolytes for the development of sodium all-solid-state batteries.

Conflicts of interest

Authors have no conflict of interest to declare.

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