Direct $^{17}$O NMR experimental evidence for Al–NBO bonds in Si-rich and highly polymerized aluminosilicate glasses†

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By using solid-state $^{17}$O NMR spectroscopy, we provide the first direct experimental evidence for bonds between Al and non-bridging oxygen (NBO) ions in aluminosilicate glasses based on rare-earth (RE) elements, where RE = (Lu, Sc, Y). The presence of ~10% Al–NBO moieties out of all NBO species holds regardless of the precise glass composition, at odds with the conventional structural view that Al–NBO bonds are absent in highly polymerized and Si-rich aluminosilicate glass networks.

Owing to their importance for both materials and earth sciences, vast efforts have been spent to improve the structural understanding of aluminosilicate (AS) glasses. Ternary M$_2$O–Al$_2$O$_3$–SiO$_2$ glasses normally involve a monovalent alkali (M$^+$) or divalent alkaline-earth (M$^{2+}$) metal cation. The glass networks comprise SiO$_4$ and AlO$_4$ tetrahedra that are corner-shared by bridging oxygen (BO) atoms. The additional negative charge of each AlO$_4$ moiety (relative to SiO$_4$) requires nearby cations for attaining local charge-balance, while the remaining M$^{2+}$ cations depolymerize the glass network by converting BO (O$^{2-}$) atoms into non-bridging oxygen (NBO; O$^{1-}$) species. The relative BO/NBO speciations in melts and glasses dictate many properties, such as viscosity, conductivity, and thermal expansion.

The following three features of the structural understanding of AS glasses have prevailed for decades, all building around the consequences of the excess negative charge of the AlO$_4$ groups: (i) Both Si and Al are four-fold coordinated (Si$^4$ and Al$^4$) by O, except if the network-modifier content is insufficient for balancing the entire Al speciation as Al$^{4+}$; then, higher-coordination AlO$_5$ and AlO$_6$ polyhedra form whenever $2n_{Al} < n_{Si}$ or $n_{Si} < n_{Al}$, where $n_E$ denotes the stoichiometric amount of element E in the glass and $z$ is the charge of M$^{z+}$. (ii) To avoid local negative charge-accumulation in the structure, there is a strong preference for Si$^{4+}$–O–Al$^{4+}$ linkages, whereas those of Al$^{4+}$–O–Al$^{4+}$ are absent (the “Loewenstein rule”). (iii) Moreover, there is a dominance of Si–NBO contacts relative to Al–NBO, implying that all NBO species are accommodated by SiO$_4$ in silica-rich AS glasses.

However, over the past decade, violations of properties (i)–(ii) are well documented for AS glasses based on mono/di-valent cations, where several studies reveal minor fractional populations of AlO$_5$ groups (a few %) in fully charge-balanced ($n_{Al} = n_{Al}/2 = n_{Si}/2$) “tectosilicate” AS glasses (notwithstanding that glasses formed at high pressure reveal significant AlO$_5$/AlO$_6$ populations). Moreover, while Loewenstein’s rule holds strictly for crystalline AS phases featuring $n_{Al} < n_{Si}$, such as zeolites and minerals, minor deviations thereof are reported for M$_2$O–Al$_2$O$_3$–SiO$_2$ glasses. Yet, whereas the early literature identified the preference of Si–NBO over Al–NBO associations, the existence of the latter were often deduced from circumstantial evidence. Nevertheless, property (iii) is nowadays assumed to apply universally for any SiO$_2$-dominated M$_2$O–Al$_2$O$_3$–SiO$_2$ glass. Direct experimental evidence for Al–NBO contacts only exist for amorphous M–Al–O aluminophase, or AS glasses that are simultaneously rich in network-modifiers and Al$_2$O$_3$, while SiO$_2$ is a minor component (<40 mol%).

However, the presence of trivalent rare-earth (RE$^{3+}$) cations in RE$_2$O$_3$–Al$_2$O$_3$–SiO$_2$ glasses leads to markedly higher configurational and chemical disorder, as mirrored in the following structural features: (1) significant AlO$_5$/AlO$_6$ populations prevail throughout the entire range of RE AS compositions, i.e., not only for those featuring $n_{Al} > n_{Si}$ and/or insufficient modifier contents. The relative amounts of higher-coordination polyhedra were demonstrated to grow for decreasing SiO$_2$ content, and particularly when the RE$^{3+}$ cation field-strength, CFS = $z/R^2$, is increased, where $R$ is the ionic radius. The markedly more cross-linked AS glass network stemming from the higher-coordination Al$^{3+}$ species was recently employed for explaining the progressively enhanced Vickers hardness observed for RE–Al–Si–O glasses with growing CFS according to La$^{3+}$ < Y$^{3+}$ < Lu$^{3+}$ < Sc$^{3+}$ < Ce$^{3+}$ < Eu$^{3+}$ (2) The high positive charge of the RE$^{3+}$ cations implies clear violations of the Loewenstein rule, reflected in a pronounced Al/Si atomic disorder for RE AS glass networks, as demonstrated by $^{29}$Si and $^{27}$Al NMR, as well as by molecular dynamics (MD) simulations. Noteworthy, the properties (1) and (2) apply generally to all AS glasses examined thus far from...
the RE = {La, Y, Lu, Sc} systems, regardless of the precise RE/Al/Si composition.10–11

Regarding the potential presence of Al\(^{p}\)-NBO contacts, i.e., violation of property (iii) of the prevailing structural picture of (Si-rich) AS glasses, we have recently presented circumstantial experimental evidence by \(^{29}\)Si NMR of a significant BO/NBO intermixing among SiO\(_4\)/AlO\(_4\) groups in La\(_2\)O\(_3\)-Al\(_2\)O\(_3\)-SiO\(_2\) structures,11 whereas MD-simulations of Y and Lu bearing glasses reveal that significant fractions (20–50\%) of all NBO species are accommodated by AlO\(_p\) groups.12 Here we provide the first direct experimental proof of significant Al-NBO contacts in SiO\(_2\)-rich RE\(_2\)O\(_3\)-Al\(_2\)O\(_3\)-SiO\(_2\) glasses with RE = {Y, Lu, Sc}, by utilizing magic-angle spinning (MAS) \(^{17}\)O NMR. Each specimen was prepared with 42–65 mol% SiO\(_2\) and \(n\) fractions (20–50\%) of all NBO species are accommodated by AlO\(_4\) groups, then referring to the peak herein assigned to Si–O[1] \(^{17}\)O-[1] species—whose peak-maximum ranges between 31–72 ppm and depending primarily on the \(n_{Al}/n_{Si}\) molar ratio—and one from NBO ions located at the SiO\(_4\) groups; Si–17O[1] (\(-137\)–158 ppm). While the \(^{17}\)O[2] NMR signal dominates, from Si–17O[1] grows concurrently with \(r\), i.e., when the glass-network polymerization decreases. Moreover, a weak but significant \(^{17}\)O resonance appears in the high-ppm region (\(-175\)–250 ppm) of all NMR spectra in Fig. 1: it is assigned to Al–17O[1] motifs. Incidentally, such a signal was previously reported by Schaller and Stebbins in the \(^{17}\)O MAS NMR spectrum from one Y\(_2\)O\(_3\)-Al\(_2\)O\(_3\)-SiO\(_2\) glass.10 However, despite noting that the “NBO peak may include oxygens bonded to AlO\(_4\) or SiO\(_4\) groups” (then referring to the peak herein assigned to Si–O[1] groups), they tentatively attributed the high-ppm signal to “NBO species with more yttrium neighbors” than those contributing to the more intense \(^{17}\)O[1] resonance. This \(^{17}\)O NMR peak appears to be a general feature of high-CFS RE-based AS glasses, but we did not detect it from La\(_2\)O\(_3\)-Al\(_2\)O\(_3\)-SiO\(_2\) glasses (data not shown), in accordance with observations made in ref. 9b.

**Table 1** RE\(_2\)O\(_3\)-Al\(_2\)O\(_3\)-SiO\(_2\) glass compositions and oxygen speciations

<table>
<thead>
<tr>
<th>Glass compositions(^{a})</th>
<th>Oxygen populations(^{b})</th>
<th>NBO populations(^{c})</th>
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<tbody>
<tr>
<td></td>
<td>(x^{[1]})</td>
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<td></td>
<td>(x^{[3]})</td>
<td>(x^{[3]})</td>
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<tr>
<td>(Y(_{28})(2.21))</td>
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<td>27.70</td>
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<td>(Sc(_{28})(2.07))</td>
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<tr>
<td>(Sc(_{28})(2.21))</td>
<td>16.96</td>
<td>28.48</td>
</tr>
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\(^{a}\) Nominal \(a\)RE\(_2\)O\(_3\)-bAl\(_2\)O\(_3\)-cSiO\(_2\) glass composition with \(a + b + c = 100 \text{ mol}\%\).\(^{b}\) MD-derived fractional populations of oxygen coordinations \(x^{[p]}\) with \(p = (0, 1, 2, 3)\), where only bonds to Si and Al are counted to define the coordination number \(p\).\(^{c}\) Fractional populations of NBO species \(x^{[1]}\) obtained by \(^{17}\)O MAS NMR, shown together with the contributions from Si–NBO \(x^{[1]}\) and Al–NBO \(x^{[1]}\) species, where \(x^{[1]} = x^{[1]} + x^{[1]}\). Values within parentheses are the corresponding MD-derived data. The uncertainties are \(\pm 0.015\) and \(\pm 0.010\) for the populations derived from NMR and MD simulations, respectively.

Note that neither the \(^{17}\)O MAS NMR spectra (Fig. 1) nor their \(^{27}\)Al counterparts may directly inform about the presence of...
Al–NBO contacts. The $^{27}$Al NMR spectra were recently reported for Y, Lu, and Sc AS glasses, all revealing coexisting AlO$_4$, AlO$_5$, and AlO$_6$ groups. Yet, unambiguous evidence for the assignment of the high-shift $^{17}$O resonance to Al–$^{17}$O$^{[1]}$ species is provided by the $^{17}$O$^{[27]}$Al TRAPDOR NMR$^{12}$ data presented in Fig. 2. Here the $^{17}$O–$^{27}$Al dipolar interaction is recoupled by applying a strong radio-frequency (rf) pulse for $\tau_{\text{rec}} = 2.5$ ms. For all $^{17}$O sites in close spatial proximity to $^{27}$Al, an attenuated integrated $^{17}$O NMR signal intensity [$S(t_{\text{rec}})$] results relative to that observed in the absence of $^{17}$O–$^{27}$Al recoupling by using a spin-echo [$S_0(t_{\text{rec}})$]. Indeed, due to the presence of Si–O–Al and Al–O–Al structural motifs, the BO-deriving $^{17}$O NMR signals manifest a significant signal dephasing. This also applies to the $^{17}$O resonance $\sim$175–250 ppm in Fig. 2 (assigned to Al–$^{17}$O$^{[1]}$ bonds), as is evidenced by its high dephasing ratio $\Delta S/S_0 = [S_0(t_{\text{rec}}) - S(t_{\text{rec}})]/S_0(t_{\text{rec}})$ obtained by deconvoluting the net $^{17}$O NMR peakshapes, as exemplified for the $S_0(t_{\text{rec}})$ spectra in Fig. 2 (b, d and f).

In contrast, the “primary” NBO-stemming resonance reveals no dephasing within the experimental/deconvolution uncertainties; the deconvolution results of Fig. 2 (b, d and f) verify that the apparent reduction of this signal stems exclusively from its overlap with the (indeed dephasing) $^{17}$O$^{[2]}$ resonances. This strongly suggests that the main $^{17}$O$^{[1]}$ NMR peak originates exclusively from Si–NBO fragments, also verifying the absence of contributions from Al–NBO moieties to this signal, as corroborated further by the additional TRAPDOR NMR data shown in Fig. 3. Fig. 3 also includes $^{17}$O NMR spectra recorded by the $^{27}$Al→$^{17}$O RAPT-CP technique. In this experiment, solely $^{17}$O species in close proximity to $^{27}$Al are detected. Indeed, while the $^{17}$O resonance-range stemming from BO structural sites is very similar to that observed directly by using central-transition (CT) selective single pulses, no significant NMR-signal intensity is observed in the region $\gtrsim$125 ppm that is primarily associated with Si–$^{17}$O$^{[1]}$ moieties (note that the weak $^{27}$Al→$^{17}$O polarization-transfer efficiency coupled with the low abundance of Al–$^{17}$O$^{[1]}$ groups precludes their observation).

Each MAS NMR spectrum of Fig. 1 was deconvoluted into signal contributions from $^{17}$O$^{[2]}$, Si–$^{17}$O$^{[1]}$, and Al–$^{17}$O$^{[1]}$ moieties (see the ESI†). The fractional populations are presented in Table 1, together with MD-derived O speciations (see ref. 9e–h). $^{17}$O NMR reveals fractional populations $x_{\text{O}}^{[1]} \approx 0.015$–0.04 of Al–O$^{[1]}$ species. An overall good agreement is observed between experiments and simulations for the total NBO population ($x^{[1]} = x_{\text{O}}^{[1]} + x_{\text{Al}}^{[1]}$), the main discrepancy being clearly over-estimated Al–NBO contacts in the glass models. The latter also reveal non-negligible populations of oxygen triclusters ($x^{[3]}$) and “free O$^{2-}$ ions” ($x^{[0]}$), as discussed further in ref. 9e–f, h. The attribution of the weak NMR signal to Al–$^{17}$O$^{[1]}$ motifs is consistent with the following trends/observations:

(i) As expected from Al–NBO fragments, there is a concomitant increase of $x_{\text{O}}^{[1]}$ with the total NBO content in the glass structure (compare the results for Lu$^{28}_{53}(2.21)$ with Lu$^{25}_{51}(2.45)$ in Table 1), as well as with the Al content (compare Lu$^{28}_{53}(2.21)$ and Lu$^{25}_{51}(2.45)$).

(ii) The isotropic chemical shifts associated with the various $^{17}$O$^{[1]}$ sites vary significantly with the nature of the RE$^{3+}$ cation, as is also evident from the NMR spectra of Fig. 1. Yet, the Al–$^{17}$O$^{[1]}$ isotropic shifts remain consistently $\sim$60–100 ppm higher than their Si–$^{17}$O$^{[1]}$ counterparts, in qualitative accordance with reported trends of Ca-based aluminate and Si-poor/Ca-rich AS glasses (we stress, however, that no Al–NBO signals were observed for Ca AS glasses exhibiting >20 mol% SiO$_2$).

(iii) The low (average) quadrupolar products $\bar{C}_{\text{O}}^{[1]} = \bar{C}_{\text{O}}(1 + \eta^{3/2}/3) \approx 2.6$ MHz observed for the Si–$^{17}$O$^{[1]}$ sites are consistent with previous reports from AS glasses, whereas the Al–$^{17}$O$^{[1]}$ species reveals $\bar{C}_{\text{O}}^{[1]} \approx 1.7$ MHz (obtained by spectra deconvolution; see the ESI†). This is to our knowledge the first estimate of quadrupolar products for Al–$^{17}$O$^{[1]}$ sites. We note that its lower value relative to Si–$^{17}$O$^{[1]}$ is expected from the higher ionic character of the Al–$^{17}$O$^{[1]}$ bond and consistent with the well-established trend observed for $^{17}$O$^{[2]}$ sites: $\bar{C}_{\text{O}}^{[2]}(\text{Al–O}) < \bar{C}_{\text{O}}^{[2]}(\text{Al–O–Si}) < \bar{C}_{\text{O}}^{[2]}(\text{Si–O–Si})$. The NMR parameters of the various $^{17}$O species will be discussed in detail elsewhere.

To conclude, we have provided the first direct experimental evidence for Al–NBO contacts in highly polymerized RE$_2$O$_3$–Al$_2$O$_3$–SiO$_2$ glasses with variable RE/Al/Si contents. The results are corroborated by MD simulations. Given the following MD-derived propensity trends of AlO$_p$ groups to associate with NBO species,
AlO₄ > AlO₅ > AlO₆,⁹₈ we attribute most of the Al-associated NBO species to be located at AlO₄ tetrahedra. Notwithstanding a strong preference for SiO₄ groups to accommodate the NBO ions, the presence of Al–NBO moieties of ≤4% out of the total O speciation (7–11% of all NBO) appears to be a general feature of AS glasses that incorporate trivalent cations with high field strength: apparently they stabilize otherwise energetically disfavoured structural motifs.

While the relative Al–NBO populations grow concurrently with the Al content of the glass, they persist in SiO₂-rich networks (at least up to ≈65 mol% SiO₂), despite that their net NBO population remains relatively low (≤4%). This is in stark contrast to AS glasses based on low-CFS mono/divalent cations, where non-negligible Al–NBO contacts have hitherto only been observed directly for fragmented networks rich in modifiers (≥50 mol% M(II)) and simultaneously featuring low SiO₂ (≤30 mol%) contents and high molar ratios n(Al₂O₃)/n(SiO₂) > 2.⁴⁴

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Notes and references


