Enhanced thermal stability of high-bismuth borate glasses by addition of iron

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Enhanced thermal stability of high-bismuth borate glasses by addition of iron

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Abstract
Glasses with nominal molar composition 20B₂O₃ – (80-x)Bi₂O₃ – xFe₂O₃ (where x = 0–40) were successfully prepared by melt-quenching. These glasses were characterised by multiple techniques including density, X-ray diffraction (XRD), X-Ray fluorescence (XRF), Raman, FT-IR and Mössbauer spectroscopies, dilatometry and differential thermal analysis (DTA). Partial replacement of Bi₂O₃ by Fe₂O₃ leads to decreasing density and molar volume and a substantial increase in thermal stability, as measured by several parameters, with maximum improvements achieved when x = 10-20. These improvements are accompanied by modest increases in dilatometric softening point. FT-IR and Raman spectra confirm the presence of BO₃ and BiO₆ structural units in all glasses, with glass structure apparently little affected by Fe₂O₃. Mössbauer spectroscopy confirms that iron is present partly as 4-fold coordinated Fe³⁺ in all glasses, with some 5- and / or 6- coordinated Fe³⁺ sites also present.

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1. Introduction
Low-melting glasses have a wide range of applications and are essential to providing hermetic sealing in many applications including electronic, semiconducting, optical and photovoltaic devices, amongst many others [1-4]. Some of the lowest-melting and lowest-sealing temperature glasses, often referred to as solder glasses, were traditionally based on the lead borate system [1-4]. Freiser [2] noted that the viscosity-temperature profile of solder glasses should be such that the “preferred zone” of viscosity enabling viscous flow of the solder glass, should leave the work piece (metal or glass) unaffected. Takamori [3] considered solder glasses
on the basis of their thermal expansion coefficient. However, in recent years legislation has effectively banned the use of lead in sealing glasses [1] and in many other glasses, and manufacturers continue to search for alternative, lead-free sealing glasses capable of delivering low sealing temperatures and acceptable thermal expansion behaviour.

Other low-melting borate glasses include alkali borates [5-8] and in order to access glass transition temperatures, $T_g$, of $< 573K$, alkali borate glasses with high alkali contents are required [6]. However, such high-alkali glasses display high thermal expansion coefficients ($15 – 25 \times 10^{-6} K^{-1}$) [6], which will render them unsuitable for some sealing applications. Lower alkali borate glasses exhibit $T_g$’s of ca. 623 - 723 K [8]. Binary alkaline earth borate glasses and ternary alkali-alkaline earth borate glasses exhibit even higher values of $T_g$, as discussed by Wozniak and James [9]. Furthermore, alkali borate glasses exhibit poor chemical durability and are therefore less attractive as potential lead-free sealing glasses. Zinc in borate sealing glasses has also received considerable attention [1-4], however, ZnO was traditionally a third component in $B_2O_3$-$PbO$ glasses and comparable high-zinc glasses in the binary $B_2O_3$-$ZnO$ system typically exhibit higher $T_g$’s $> 773 K$, too high for some sealing applications.

One family of glasses that may be capable of replacing lead with a non-toxic alternative in some applications occurs in the $B_2O_3$-$Bi_2O_3$ system. In his recent, very thorough review of bismuth-containing glasses Maeder [1] noted that $Bi_2O_3$ “appears a quite promising drop-in replacement for PbO” but “the somewhat lower fluxing ability…..leads to higher processing temperatures”. Despite this, bismuth-rich low temperature sealing glasses have been successfully developed [10-12]. Bismuth borates have been stated to form transparent glasses over a particularly wide range of $Bi_2O_3$ contents, from 20 to 85 mol% [13]. Bajaj et al. [13] undertook a detailed investigation of the structure and properties of glasses in the system $xBi_2O_3 – (100-x)B_2O_3$ where $x = 20$ to 66 (mol %), and noted the interest in this system for non-linear optics. Shaaban et al. [14] studied glasses with compositions $xBi_2O_3 – (100 – x)B_2O_3$ where $x = 35$ to 60 mol%. They carefully characterised thermal properties, crystallisation behaviour and structure, demonstrating that thermal stability decreased with increasing $Bi_2O_3$ content.

A wide range of additional components to the $B_2O_3$-$Bi_2O_3$ glass system has been studied previously, as summarised by Maeder [1]. However, a dopant that has received relatively little attention, particularly in the context of high-bismuth sealing glasses, is iron. Dumbaugh [15] showed that the binary system $Bi_2O_3$-$Fe_2O_3$, which contains no classical glass formers, is capable
of forming glasses over a considerable range of iron contents, given the inclusion of a third component (in Dumbaugh’s case CdO and / or PbO). This offered a tantalising clue to the potential for incorporating iron in high-bismuth glasses. Qiu et al. [16] studied glass formation and DC conductivity in the system B₂O₃-Bi₂O₃-Fe₂O₃, and confirmed that high-Bi₂O₃, low-B₂O₃ glasses containing up to 40 mol % Fe₂O₃ can be formed by press-quenching the melt between two copper blocks. Baia et al. [17, 18] studied the structure of a wide range of compositions in the systems (mol%) 95[xB₂O₃,(1-x)Bi₂O₃]-5Fe₂O₃ where 0.07 ≤ x ≤ 90 and 90[xB₂O₃,(1-x)Bi₂O₃]-10Fe₂O₃ where 0.07 ≤ x ≤ 0.625 using Raman and FT-IR spectroscopies. They found that Bi³⁺ cations are incorporated in the glass network as BiO₆ polyhedra, and iron doping can stabilise the glass network at intermediate bismuth contents. El-Desoky and colleagues [19-21] studied doping of iron into B₂O₃-Bi₂O₃-R₂O (R=Li, Na, K) glasses, however, they were primarily investigating electrical properties and their glasses were relatively high in B₂O₃ and low in Bi₂O₃. Akamatsu et al. [22] studied the magnetic and structural properties of glasses with nominal molar composition xFe₂O₃ - (80 − x) Bi₂O₃ - 20 B₂O₃ (where x = 18.2 - 40.0). These glasses exhibit unusual magnetic behaviour which they explained in terms of coexisting spin glass phases and magnetic clusters. Another application area of interest in high-bismuth glasses is for radiation shielding / dosimeter applications, and these have also received recent attention [23 - 24]. The presence of boron and bismuth could enable potential applications in the nuclear arena, due to the high absorption cross-section for thermal-neutrons (B) and γ-radiation (Bi). In terms of iron-free bismuth borate glasses, Yawale and Pakade [25] studied the physical and electrical transport properties of Bi₂O₃-B₂O₃ glasses, and further detailed studies of the physical properties of Bi₂O₃-B₂O₃ glasses, across a wide range of compositions, were carried out by Stehle et al. [26] and George et al. [27]. The high densities of bismuth-rich borate glasses are inherently linked to all of the above applications. As noted by Maeder [1], anomalies in density and T_g can also suggest structural information, hence knowledge of the density and molar volume of bismuth borate glasses is important. One additional property requirement of sealing glasses is that they exhibit sufficient chemical durability for the application in question. As discussed by Maeder [1], some high-lead borate glasses and high-bismuth borate glasses exhibit chemical durabilities that are considered acceptable for many low-temperature sealing applications.
Understanding the effects of iron additions to the structure and properties of high-bismuth borate glasses is therefore of interest, not only from the perspective of developing new lead-free sealing glasses, but also within the wider context of understanding the structure of high-bismuth oxide glasses, and in terms of new potential applications requiring novel electrical, nuclear or other properties. Here we have studied the effects of iron doping on the structure and physical properties of glasses with nominal molar composition 20 B₂O₃ – (80-x) Bi₂O₃ – xFe₂O₃ (where x = 0–40).

2. Experimental Procedures

Batches to provide 50g of glass were produced using the reagent-grade chemicals boric acid (H₃BO₃, 99.5%), bismuth oxide (Bi₂O₃, 99%) and iron oxide (Fe₂O₃, 98%). Raw materials were carefully dried overnight prior to weighing (H₃BO₃ at 323 K so as not to decompose; and Bi₂O₃ and Fe₂O₃ at 393 K). Batches were weighed out using a calibrated 2 decimal-place balance, and were then mixed thoroughly. Nominal glass compositions are listed in Table 1. The mixed batches were then placed in recrystallised Al₂O₃ crucibles, with an Al₂O₃ lid placed over the crucible to reduce volatilisation losses during heating and melting. Crucibles were placed in an electric furnace and heated at 5 K / min in air to 1373 K, and held at this temperature for 30 minutes. The lids were then removed from the crucibles and the crucibles removed from the furnace and their contents press-quenched between two steel plates. Resulting samples were then stored in sealed bags. Samples were powdered in an attrition mill for 1 minute to provide samples for XRD, XRF, Mössbauer spectroscopy, DTA and FT-IR spectroscopies. Density measurements, Raman spectroscopy and dilatometry were performed on bulk glass samples.

Densities have been measured by the Archimedes method using deionised water as the suspension medium. Archimedes densities are calculated using (1):

\[
\text{Density} = \left(\frac{W_A}{W_A - W_W}\right) \times \delta_W \quad (1)
\]

where \(W_A\) = weight in air, \(W_W\) = weight in water and \(\delta_W\) = temperature correction. Averages of three measurements were taken for each sample. Molar volumes have been calculated using measured densities and analyzed compositions.
Elemental analysis was performed using a PANalytical MagiX Pro XRF spectrometer equipped with a Rh anode. Powdered samples were mixed with cellulose binder and pressed into pellets using a 20 tonne force in a Retsch PP40 hydraulic press. A semi-quantitative XRF analysis program, IQ+ for standardless data analysis, was used here. Boron was not detected by XRF and so to enable useful analysis it had to be assumed that no boron losses had occurred during melting, and analysed content was modified to take account of the presence of boron on a pro rata basis. Accuracies of the resultant data in this study, arising from the use of the semi-quantitative analysis program, are estimated to be ±3 of their values as provided by the analysis, based on historical results using this instrument and analysis program on other, related glasses.

X-ray diffraction was performed at room temperature on powdered pellets using an Empyrean PANalytical diffractometer in Bragg–Brentano geometry. Samples were mounted on a reflection/transmission spinner stage rotating at 15 rpm, irradiated with Cu Kα radiation (λ = 1.5406 Å) over a 2θ range of 20–80°, with step size 0.013 2θ and step time 68.59 s, 10 mm incident mask, 1° antiscatter and divergence slits, and a nickel beta filter. X-rays were detected using a PIXCEL-3D area detector.

Dual DTA and TGA analysis of the sample were carried using a NETZSCH STA 409 PC/PG, with 180mg of powdered sample in air. Measurements were carried out at a heating rate of 10°C / minute from 50°C to 780°C. No detectable changes in sample mass during analysis were noted, within the uncertainties of the instrument. The temperatures \( T_g \) (glass transition temperature), \( T_x \) (onset crystallization temperature) and \( T_c \) (crystallization peak temperature) were determined from the corrected DTA traces, which are shown in Figure 2. The extracted \( T_g \), \( T_x \) and \( T_c \) temperatures are given in Table 1. Uncertainties associated with each of the above measured parameters (+/- 3K) were estimated from repeat measurements of the same sample and temperature calibration of the instrument using a standard. Thermal stabilities and glass-forming abilities of the glasses were estimated using three different methods: \( \Delta T = (T_x - T_g) \) [28]; \( S = (T_c - T_x) (T_x - T_g)/(T_g) \) [29]; and the Hrubý parameter, \( K_{gl} = (T_x - T_g) / (T_m - T_x) \) [30]. Figure 3 shows \( T_g \) and \( \Delta T = (T_x - T_g) \) as functions of nominal Fe\(_2\)O\(_3\) content.

Dilatometry was carried out using a Netzsch Expedis Select dilatometer at a heating rate of 5 K / min from 293 K to a temperature above the dilatometric softening point, \( T_d \). Owing to the press-quenching method used to produce the glass samples and their resulting thickness of ~1 mm, it was only possible to obtain \( T_d \) values from dilatometry and not coefficient of thermal
expansion, $\alpha$. The obtained $T_d$ values are shown in Table 1. Uncertainties associated with measured $T_d$ (+/- 3K) were estimated from repeat measurements of the same sample and temperature calibration of the instrument using a standard.

Fourier-Transform Infra-Red (FT-IR) spectroscopy was carried out using a Nicolat Nexus 470 FT-IR spectrometer. Powdered glass samples were mixed with dry KBr and pressed into pellets using a pellet press, for transmission measurements. Spectra were recorded between 600 and 1800 cm$^{-1}$, and are shown in Figure 4.

Raman spectroscopy was carried out using a Thermo Scientific DXR2 spectrometer with a depolarised 10 mW 532 nm laser, on bulk samples of all glasses, between 200 and 2000 cm$^{-1}$. Calibrations with the proprietary Thermo alignment tool were carried out before, during and after each sample measurement. Spectra were baseline-corrected and are shown in Figure 5 and key Raman band assignments associated with the spectral features are summarised in Table 3. There is a strong damping effect, such that most spectral features were increasingly suppressed upon increasing Fe$_2$O$_3$ content of the glasses. The spectra shown in Figure 5 have thus been progressively amplified in order to illustrate spectral features, which explains the progressively lower signal-to-noise (S/N) ratio with increasing Fe$_2$O$_3$ content of the samples.

Transmission $^{57}$Fe Mössbauer spectroscopy measurements were carried out using acrylic absorber discs with a sample area of 1.767 cm$^2$. These were loaded with sample to present $2.16 \times 10^{-3}$ g cm$^{-2}$ of Fe to achieve a Mössbauer thickness of 1. Sample weights of 0.013 g were homogeneously mixed with graphite to achieve this level of loading. The 14.4 keV $\gamma$-rays were supplied by the cascade decay of 25 mCi $^{57}$Co in Rh matrix source, oscillated at constant acceleration by a SeeCo W304 drive unit, and detected using a SeeCo 45431 Kr proportional counter operating with 1.745 kV bias voltage applied to the cathode. All measurements were carried out at room temperature over a velocity range of $\pm$4 mm s$^{-1}$, and were calibrated relative to $\alpha$-Fe foil. Spectral data were fitted using the Recoil software package [31], using Lorentzian line shapes. The low S/N obtained for each sample was caused by strong absorption of incident gamma-rays by the sample (due to very high bismuth contents). Long collection times of up to 3 weeks were required for each sample, and consequently spectral fitting was limited to a single doublet for each spectrum. Attempts were made to fit two doublets, however, these resulted in no significant improvement in reduced $\chi^2$ for each fit and were consequently discarded in favour of single-doublet fits.
3. Results

Table 1 shows the nominal and analysed glass compositions studied, with sample name nomenclature (0Fe, 10Fe, 20Fe, 30Fe, 40Fe) reflecting the nominal molar percentage Fe$_2$O$_3$ content. X-ray diffraction confirmed the X-ray amorphous nature of all samples. Standardless XRF analysis confirms that all of our glasses contain ~1 to 2 wt% (~4 to 7 mol%) Al$_2$O$_3$, as shown in Table 1. These Al$_2$O$_3$ contents are attributed to unavoidable chemical interactions of the molten glass with the Al$_2$O$_3$ crucibles during melting. Due to the very high Bi$_2$O$_3$ contents of these glasses (which strongly re-absorb emitted X-rays), combined with the semi-quantitative nature of the standardless XRF analysis program used, there are considerable uncertainties associated with the XRF analyses, as shown in Table 1. Measured densities and corresponding molar volumes, shown in Table 1 and Figure 1, depict monotonic decreases and increases, respectively, in density and molar volume with increasing Fe$_2$O$_3$ content.
Table 1. Nominal (analysed) glass compositions, density, molar volume and thermal parameters.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Nominal (analysed) composition/mol%</th>
<th>Density, ρ/g cm⁻³ (± 0.01)</th>
<th>Molar Vol., Vₘ cm⁻³ (± 0.5)</th>
<th>Tᵥ/K (± 3)</th>
<th>Tₓ/K (± 3)</th>
<th>T𝑐/K (± 3)</th>
<th>Tₘ/K (± 3)</th>
<th>ΔT/K (±6) [28]</th>
<th>S [29]</th>
<th>K₉ [30]</th>
<th>T₉/K (±3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0Fe</td>
<td>20.0* 0.0 (0.0) 80.0 (73.7) 0.0 (6.3)</td>
<td>7.88</td>
<td>49.1</td>
<td>603</td>
<td>635</td>
<td>655</td>
<td>861</td>
<td>32</td>
<td>1.06</td>
<td>0.142</td>
<td>600</td>
</tr>
<tr>
<td>10Fe</td>
<td>20.0* 10.0 (6.8) 70.0 (65.9) 0.0 (7.3)</td>
<td>7.58</td>
<td>47.0</td>
<td>635</td>
<td>701</td>
<td>715</td>
<td>858</td>
<td>66</td>
<td>1.46</td>
<td>0.420</td>
<td>635</td>
</tr>
<tr>
<td>20Fe</td>
<td>20.0* 20.0 (14.5) 60.0 (61.0) 0.0 (4.5)</td>
<td>7.25</td>
<td>44.9</td>
<td>662</td>
<td>723</td>
<td>755</td>
<td>923⁴</td>
<td>61</td>
<td>2.95</td>
<td>0.305⁴</td>
<td>665</td>
</tr>
<tr>
<td>30Fe</td>
<td>20.0* 30.0 (21.5) 50.0 (52.7) 0.0 (5.7)</td>
<td>6.99</td>
<td>42.2</td>
<td>692</td>
<td>731</td>
<td>752⁷</td>
<td>948³</td>
<td>39</td>
<td>1.18</td>
<td>0.180³</td>
<td>675</td>
</tr>
<tr>
<td>40Fe</td>
<td>20.0* 40.0⁴ 40.0⁴ 0.0⁴</td>
<td>6.63</td>
<td>39.9</td>
<td>690</td>
<td>745</td>
<td>777⁷</td>
<td>904⁴</td>
<td>55</td>
<td>2.55</td>
<td>0.346⁴</td>
<td>680</td>
</tr>
</tbody>
</table>

* not measured; ⁴ DTA traces showed multiple phase formation / melting.
Figure 1. Measured densities and estimated molar volumes for samples 0Fe to 40Fe. Lines are shown as a guide. Error bars are smaller than data points.

Differential Thermal Analysis (DTA) traces, shown in Figure 2, and the extracted and calculated thermal parameters given in Table 1, depict clear trends in $T_g$, the glass transition temperature; in $T_x$, the onset crystallisation temperature, and in $T_c$, the peak crystallisation temperature, for this series of glasses. $T_g$, $T_x$ and $T_c$ all increase with increasing nominal Fe$_2$O$_3$ content, but $\Delta T$, the temperature difference between $T_x$ and $T_g$ ($\Delta T = T_x - T_g$), varies with glass composition. This behaviour is tabulated in Table 1 and shown graphically in Figure 3 for $T_g$ and the temperature difference between $T_x$ and $T_g$ ($\Delta T = T_x - T_g$). Figure 2 also shows a clear endothermic event, associated with melting, for the 0, 10 and 20 mol% Fe$_2$O$_3$ glasses. This endotherm moves to slightly lower temperatures with iron additions, from ca. 888 K at 0 mol % Fe$_2$O$_3$ to ca. 863 K at 20 mol% Fe$_2$O$_3$. At higher Fe$_2$O$_3$ additions, this endothermic event disappears and with increasing iron content the high-temperature region of the DTA traces becomes increasingly complex. In addition to thermal stability related factors, high temperature viscosity (as measured by dilatometric softening point, $T_d$) shows a monotonic increase from
600K at 0 mol% Fe$_2$O$_3$ to 635K at 10 mol% Fe$_2$O$_3$ to 665K at 20 mol% Fe$_2$O$_3$. At higher Fe$_2$O$_3$ the rate of increase in $T_d$ decreases, with $T_d$ of 675K at 30 mol% Fe$_2$O$_3$ and 680K at 40 mol% Fe$_2$O$_3$.

*Figure 2.* DTA traces for samples 0Fe to 40Fe
Figure 3. Onset $T_g$ and thermal stability ($T_x - T_g$) as functions of nominal Fe$_2$O$_3$ content

Fourier-Transform Infra-Red (FT-IR) transmission spectra, shown in Figure 4, evidence little change throughout the series of glasses. Absorption bands are as follows: (i) a narrow band centred at 720 cm$^{-1}$; (ii) a broad, weak band centred at 920 cm$^{-1}$; (iii) a strong, broad band centred at 1180 cm$^{-1}$; and (iv) a weaker shoulder at 1280 cm$^{-1}$. Raman spectra for all samples (Figure 5) all exhibit the following bands: (i) a broad band centred at 400 cm$^{-1}$; (ii) a broad band at 600 cm$^{-1}$; (iii) a weak, narrow band at 710 cm$^{-1}$; (iv) a sharp band at 840 cm$^{-1}$; (v) a weak band at 920 cm$^{-1}$; and (vi) a broad band centred at 1230 cm$^{-1}$ comprising two overlapping bands centred at 1180 cm$^{-1}$ and 1280 cm$^{-1}$. Table 2 summarises FT-IR and Raman band assignments and provides supporting references.

Mössbauer spectra (Figure 6) and accompanying fitted parameters (Table 3) show closely similar Centre Shift (CS), Quadrupole Splitting (QS) and Linewidth (LW) values for all three samples studied, containing nominally 20, 30 and 40 mol% Fe$_2$O$_3$, indicating little or no change in the oxidation state or local environment of Fe cations with changing glass composition.
Figure 4. FT-IR spectra for samples 0Fe to 40Fe

Figure 5. Raman spectra for samples 0Fe to 40Fe (amplified with increasing Fe content)
Table 2. IR and Raman band assignments

<table>
<thead>
<tr>
<th>IR / cm⁻¹</th>
<th>Raman / cm⁻¹</th>
<th>Structural origin</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>400</td>
<td>Bi-O-Bi + Bi-O in [BiO₆] units.</td>
<td>[17, 18, 23, 32, 33]</td>
</tr>
<tr>
<td>-</td>
<td>620</td>
<td>Bi-O (NBO) stretching vibration</td>
<td>[17, 18, 33]</td>
</tr>
<tr>
<td>720</td>
<td>720</td>
<td>B-O-B bending in trigonal BO₃ units</td>
<td>[18, 32, 33, 34]</td>
</tr>
<tr>
<td>-</td>
<td>840</td>
<td>Bi-O in pyramidal (BiO₃) units and B-O in (BO₄) units</td>
<td>[18, 32, 33]</td>
</tr>
<tr>
<td>880</td>
<td></td>
<td>Bi-O / Bi-O-Bi in BiO₆ octahedra</td>
<td>[18]</td>
</tr>
<tr>
<td>920</td>
<td>920</td>
<td>B-O in trigonal (BO₃) units</td>
<td>[18]</td>
</tr>
<tr>
<td>900-950</td>
<td>-</td>
<td>Stretching vibration of (BO₄) units</td>
<td>[32, 33, 35]</td>
</tr>
<tr>
<td>1150-1170</td>
<td>1180</td>
<td>B-O in tetrahedral (BO₄) units</td>
<td>[18, 34]</td>
</tr>
<tr>
<td>1200-1300</td>
<td>1280</td>
<td>B-O of (BO₃) triangles</td>
<td>[18, 23, 32, 34, 36]</td>
</tr>
</tbody>
</table>

Table 3. Fitted Mössbauer parameters for samples 20Fe to 40Fe

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fitting Reduced $\chi^2$</th>
<th>Centre Shift ± 0.02 / mm s⁻¹</th>
<th>Quadrupole Splitting ± 0.02 / mm s⁻¹</th>
<th>Linewidth HWHM ± 0.02 / mm s⁻¹</th>
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</thead>
<tbody>
<tr>
<td>20Fe</td>
<td>0.517</td>
<td>0.31</td>
<td>0.88</td>
<td>0.26</td>
</tr>
<tr>
<td>30Fe</td>
<td>0.621</td>
<td>0.34</td>
<td>0.86</td>
<td>0.27</td>
</tr>
<tr>
<td>40Fe</td>
<td>0.493</td>
<td>0.32</td>
<td>0.90</td>
<td>0.23</td>
</tr>
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</table>
Figure 6. Fitted Mössbauer spectra for samples 20Fe to 40Fe
4. Discussion

Other researchers [25-27] have studied the same nominal base glass composition as ours (80 mol% Bi₂O₃ – 20 mol% B₂O₃). Yawale and Pakade [25] melted their glasses at 1273 - 1773 K for 2 hours in fireclay (aluminosilicate) crucibles and obtained a density of 7.701 g cm⁻³. On the other hand, Stehle et al. [26] melted their glass at considerably lower temperatures and times (1273 K for 10 minutes) in platinum crucibles and obtained a density of ~8.5 ± 0.2 g cm⁻³. Our measured density of 7.88 ± 0.01 g cm⁻³ for glass melted in an Al₂O₃ crucible at 1373 K for 30 min lies between these literature data. The differences in measured densities are likely to be principally due to different levels of contamination from corrosion of the crucible material (negligible for platinum [26, 27]; contamination from Al₂O₃ (this study) and fireclay [25]). There may also be contributions from different levels of volatilisation loss on account of differences in melting time and temperature (lowest for 1273 K / 10 minutes [26]; slightly higher for 1373 K / 30 minutes (this study); higher for 1273 - 1773 K for 10 minutes [27]; and considerably higher for 1273 - 1773 K for 2 hours [25]. Whilst none of these other authors [25-27] chemically analysed their glasses, Stehle et al. [26] did carry out “weight loss measurements to ensure stoichiometric accuracy”. It is thus reasonable to assume that their glasses exhibit as close to the true density of glasses with the composition 80 mol% Bi₂O₃ – 20 mol% B₂O₃, free from contamination, and produced under the roller quenching preparation conditions they used, as can be achieved experimentally. All of our glasses were found to contain ~1 to 2 wt% (~4 to 7 mol%) Al₂O₃, as detected by standardless XRF analysis and shown in Table 1. The limited accuracy of XRF IQ+ program makes it difficult to gauge boron losses, as may reasonably be expected to occur at some level, through consideration of the contents of the other main components, Bi₂O₃ and Fe₂O₃. However, we note that whilst the analysed Bi₂O₃ contents are lower than their nominal values for samples 0Fe and 10Fe, they are above their nominal values for samples 20Fe and 30Fe. Consequently, and also taking into account the analysed Al₂O₃ contents of the glasses which will reduce the contents of all other components proportionately, we can conclude that boron losses, whilst not necessarily negligible, were not high.

Thermal analysis results clearly show that modest replacement of Bi₂O₃ by Fe₂O₃ (10-20 mol %) results in a significant expansion of the temperature range between Tₘ and Tₓ, from 32 K in the iron-free glass which essentially doubles to 66 K in the 10 mol % Fe₂O₃ glass. Further
replacements of Bi$_2$O$_3$ by Fe$_2$O$_3$ (20-40 mol%) result in decreases in $\Delta T$ from this maximum – however, even at 40 mol% Fe$_2$O$_3$, the value of $\Delta T$ is still larger than for the iron-free 20 B$_2$O$_3$ – 80 Bi$_2$O$_3$ glass. The increasing suppression of the endothermic event observed upon increasing additions of iron suggest frustration of crystallisation, producing multiple phases, as evidenced by the multiple weak crystallisation / melting peaks at temperatures above ca. 823 K in the DTA traces for the iron-containing samples. The extracted thermal stability and glass formation criteria $\Delta T$, S and $K_{gl}$ are all consistent with the view that partial replacement of Bi$_2$O$_3$ by Fe$_2$O$_3$, especially at 10 and 20 mol % replacement, enhanced glass formation and thermal stability. Comparison of extracted thermal properties $\Delta T$, S and $K_{gl}$ with data for similar glasses from literature shows that the obtained thermal parameters for the 80 Bi$_2$O$_3$ – 20 B$_2$O$_3$ base glass (Sample 0Fe) are consistent with the trends observed by Shaaban et al. [14] for glasses with compositions $x$Bi$_2$O$_3$ – (100 – $x$)B$_2$O$_3$ with $x$ = 35 to 60 mol%; and with Cheng et al. [32] for glasses with compositions $x$Bi$_2$O$_3$ – (100 – $x$)B$_2$O$_3$ with $x$ = 30 to 60 mol%. Increasing Bi$_2$O$_3$ content in all series lead to decreased $\Delta T$ and $K_{gl}$, and similar values of S, indicating a trend of decreasing glass stability. For example, Shaaban’s 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass gave $\Delta T$ = 56K, S = 0.954 and $K_{gl}$ = 0.386; and Cheng’s 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass gave $\Delta T$ = 68K and $K_{gl}$ = 0.42 whilst our 80 Bi$_2$O$_3$ – 20 B$_2$O$_3$ glasses give $\Delta T$ = 32K, S = 1.061 and $K_{gl}$ = 0.142, confirming the trend in decreasing glass stability with increasing Bi$_2$O$_3$ content and decreasing B$_2$O$_3$ content, which is consistent with Stehle et al. [26]. However, in our samples, the introduction of Fe$_2$O$_3$ leads to a rapid increase in glass stability wherein the composition 70 Bi$_2$O$_3$ – 20 B$_2$O$_3$ – 10 Fe$_2$O$_3$ gave $\Delta T$ = 66K, S = 1.455 and $K_{gl}$ = 0.420, which is more stable than Shaaban’s 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass and has the same stability as Cheng’s 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass. Our 60 Bi$_2$O$_3$ – 20 B$_2$O$_3$ – 20 Fe$_2$O$_3$ is also more stable than Shaaban’s 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass, with $\Delta T$ = 61K, S = 2.948 and $K_{gl}$ = 0.305. Study of the data of Stehle et al. [26] for binary Bi$_2$O$_3$-B$_2$O$_3$ glasses enables further comparison of the effect of Fe$_2$O$_3$ addition vs. B$_2$O$_3$ addition. Stehle et al. [26] illustrated the difference between “the crystallisation temperature $T_x$” and onset $T_g$. In the absence of any description by Stehle et al. [26] as to whether their “$T_x$” refers to onset crystallisation temperature ($T_x$ in this study) or crystallisation peak temperature ($T_c$ in this study), we have assumed, by comparison with our data, that their “crystallisation temperature $T_x$” is the crystallisation peak temperature ($T_c$ in this study). Based on this assumption, Stehle’s $T_c$-$T_g$ increased from ~58 K for 80 Bi$_2$O$_3$ – 20 B$_2$O$_3$ glass to ~70 K for 70 Bi$_2$O$_3$ – 30 B$_2$O$_3$ glass, to
~115 K for 60 Bi$_2$O$_3$ – 40 B$_2$O$_3$ glass. By comparison our glasses with nominal 80, 70 and 60 mol% Bi$_2$O$_3$ contents exhibited T$_c$ - T$_x$ of 52, 80 and 93 K, which also mirror this trend of increasing (T$_x$-T$_g$) with decreasing Bi$_2$O$_3$ content, and are of similar magnitude. This in turn indicates that the improvements in (T$_x$-T$_g$) obtained by partially replacing Bi$_2$O$_3$ by B$_2$O$_3$ are mirrored by the improvements in (T$_x$-T$_g$) resulting from partially replacing Bi$_2$O$_3$ by (Fe$_2$O$_3$ + Al$_2$O$_3$). The effects of Al$_2$O$_3$, in addition to Fe$_2$O$_3$, must be considered for our glasses, since XRF analysis confirmed the presence of ~ 4 to 7 (± 3) mol % Al$_2$O$_3$. Previously it was shown that similar additions (5 mol%) of Al$_2$O$_3$ can improve the thermal stability of iron phosphate glasses [37]. In addition to this, increasing the number of different elements in a glass is well known to frustrate crystallisation. It is therefore likely that the levels of Al$_2$O$_3$ present in our glasses have an impact on thermal stability. A substantial number of researchers describing similar glasses also prepared their glasses in refractory crucibles (or did not state the crucible material) – and therefore it is suggested that other such glasses may have also contained similar, or greater, impurity levels of Al$_2$O$_3$ [14, 16, 17, 18, 21, 23]. Since Al$_2$O$_3$ contents of all glasses studied here remain at broadly similar levels, including the 80 Bi$_2$O$_3$-20 B$_2$O$_3$ base glass, it can be concluded that the improvements in thermal stability, compared with the base glass, that are observed in this study are as a result of partially replacing Bi$_2$O$_3$ by Fe$_2$O$_3$ and not Al$_2$O$_3$ contamination. The increases in T$_d$ with increasing Fe$_2$O$_3$ content are consistent with partial replacement of Bi$^{3+}$ by Fe$^{3+}$, as the corresponding dissociation energies of Fe$_2$O$_3$ (287 kcal / mol) and Bi$_2$O$_3$ (147 kcal / mol) [38] are consistent with a more rigid glass network requiring higher temperatures to initiate viscous flow, as Fe$_2$O$_3$ increasingly replaces Bi$_2$O$_3$.

The eutectic temperature in the Bi$_2$O$_3$-2Bi$_2$O$_3$.B$_2$O$_3$ subsystem, in which the 80 mol% Bi$_2$O$_3$ – 20 mol% B$_2$O$_3$ (sample 0Fe) composition is located, has a T$_m$ of 622°C (895 K) [39, 40]. According to a more recent version of the Bi$_2$O$_3$-B$_2$O$_3$ phase diagram [41] the 80 mol% Bi$_2$O$_3$ – 20 mol% B$_2$O$_3$ composition is very close to the eutectic composition of the Bi$_2$O$_3$-5:3Bi$_2$O$_3$.B$_2$O$_3$ subsystem. Our T$_m$ of 588°C found in this work for sample 0Fe has an intermediate value, although the glass is actually ternary, since it contains Al$_2$O$_3$, which may explain the observed, modest difference in T$_m$ from the previous work. The other compositions studied here are quaternary as both Fe$_2$O$_3$ and Al$_2$O$_3$ must be considered.

When considering the structure of the B$_2$O$_3$.Bi$_2$O$_3$.Fe$_2$O$_3$.Al$_2$O$_3$ glasses studied here, it is important to first consider the structure of binary B$_2$O$_3$.Bi$_2$O$_3$. Maeder [1] summarised a
number of studies of high-bismuth borate glasses, and in particular, Bajaj et al. [13] and Dimitrov and Komatsu [42], using $^{11}$B MAS-NMR, provided evidence strongly suggesting that over 80% of the boron in very-high lead and bismuth borate glasses (i.e. in all glasses studied here) will adopt trigonal coordination; and consequently the amount of four-coordinated B$^{3+}$ is expected to be small. Bismuth in Bi$_2$O$_3$–B$_2$O$_3$ glasses has been described as occupying (BiO$_6$) octahedral sites [1, 17, 18, 23, 32, 33]. All Raman spectra (Figure 5) show bands due to bridged anion modes at ca. 400 cm$^{-1}$. The Raman band at ca. 600 cm$^{-1}$ is attributed to Bi-O$^-$ stretching (vibration of bismuth - non-bridging oxygen) in (BiO$_6$) polyhedra [17, 18, 33]. The IR band at 720 cm$^{-1}$ is attributed to bending vibrations of B–O–B trigonal units [18, 32, 33, 34]. The weak IR band at 920 cm$^{-1}$ has been attributed to stretching vibrations of (BO$_4$) units [32, 33, 35]. The IR and Raman bands centred at 1180 cm$^{-1}$ and 1280 cm$^{-1}$ are a matter of debate, as they have received little prior attention as discrete units in the context of very high -PbO or -Bi$_2$O$_3$ borate glasses. However, it is clear that Raman bands at high frequency (ca. 1250 cm$^{-1}$) provide information on short range order in these borate glass systems.

Aside from the overall intensity of Raman-active bands being suppressed by Fe$_2$O$_3$ additions, the spectral changes that do occur upon increasing Fe$_2$O$_3$ addition, in both FT-IR and Raman spectra, are surprisingly subtle, with no large changes. Overlaying the FT-IR spectra on each other reveals little difference between any of the sample spectra. Overlaying the Raman spectra, when amplified sufficiently to counteract the suppressing effect of Fe$_2$O$_3$ additions (as in Figure 5), shows only the band at 720 cm$^{-1}$ being incorporated into the stronger background occurring at low Raman shifts, and a small change in the relative intensities of the two overlapping Raman bands at 1180 cm$^{-1}$ and 1280 cm$^{-1}$, which manifests as a change in the overall peak profile in this spectral region.

The splitting of both the high-frequency IR band and the high-Raman shift band into bands at 1180 and 1280 cm$^{-1}$ can be observed in the data of several authors [14, 17, 18, 33, 34, 35, 36, 43, 44]. However, few have specifically discussed this band splitting. It was discussed by Kotkova et al. [43], who also observed band splitting for PbO-Bi$_2$O$_3$-B$_2$O$_3$ glasses with high (Pb+Bi) contents. They noted that the band at 1180 cm$^{-1}$ is often stated to indicate of the presence diborate groups, but that it is unlikely that diborate groups form at such high modifier contents. They attributed both Raman bands, at 1180 cm$^{-1}$ and 1280 cm$^{-1}$ to trigonal borate groups. However, others have attributed the band at 1180 cm$^{-1}$ to tetrahedral borate groups.
Assuming this is the correct assignment in the case of our glasses, then our results suggest a modest increase in the $[^3]B^{3+} / [^4]B^{3+}$ ratio with increasing Fe$_2$O$_3$ content. However, as previously noted, evidence from other studies indicates that the large majority (> 80%) of the boron in our glasses is likely to be trigonally-coordinated $[^3]B^{3+}$, so further research would be required to confirm the abundance of tetrahedral $[^4]B^{3+}$ and its compositional dependence, and to relate this back to the IR and Raman spectra. On balance, the majority of literature suggests that both the 1180 cm$^{-1}$ and 1280 cm$^{-1}$ IR and Raman bands are related to trigonal $[^3]B^{3+}$.

The suppressing effect of Fe$_2$O$_3$ concentration on Raman spectra that we have observed here are also mirrored in other, related studies. Kotkova et al. [43] showed similar suppression with increasing PbO concentration in (PbO)$_x$(Bi$_2$O$_3$)$_{0.2}$(B$_2$O$_3$)$_{0.8-x}$ glasses and Ardelean and Cora [33] showed comparable behaviour with increasing CuO concentration in xCuO.(100-x)(3Bi$_2$O$_3$-B$_2$O$_3$) glasses. Research by Pan et al. [45], who studied a wide range of B$_2$O$_3$-PbO glasses, showed that the Raman scattering cross-section increased strongly with increasing PbO content, with this increase dominating in the low-frequency region. This behaviour was attributed in part to the polarizability of Pb$^{2+}$ cations. Since Bi$^{3+}$ is also highly polarisable, one possible cause of the suppression of Raman spectra upon increasing Fe$_2$O$_3$ content in our glasses is that the Fe$_2$O$_3$ partially replaced Bi$_2$O$_3$ and consequently the Raman cross-section of the glass became considerably weaker.

The Centre Shift (CS) and Quadrupole Splitting (QS) values obtained from fitting of Mössbauer spectra are consistent with the Fe$^{3+}$ oxidation state in oxide glasses [47-53] and there is no evidence for any measurable levels of Fe$^{2+}$ in any of the glasses studied here. The coordination of the Fe$^{3+}$ ions identified can be determined qualitatively from CS and QS values. With CS of 0.31-0.34 mm s$^{-1}$ and QS of 0.86-0.90 mm s$^{-1}$ these values lie on the tetrahedral side of, but very close to, the boundary between 4-coordination and higher coordination numbers of 5 and 6 [47-50]. The associated Linewidth (LW) values of 0.23-0.27 mm s$^{-1}$ are consistent with a range of site distortions, as occur in oxide glasses. It is therefore highly likely that whilst some Fe$^{3+}$ adopts tetrahedral coordination with respect to oxygen, a substantial fraction of Fe$^{3+}$ occupies 5- and / or 6-coordinated sites in these glasses: this is consistent with literature for a wide range of borate glasses [47-53]. The fact that the obtained Mössbauer CS, QS and LW parameters do not change significantly between 20%, 30% and 40% Fe$_2$O$_3$ nominal content of the glasses studied, suggests that the Fe$^{3+}$ local environments, range of site occupations and
range of site distortions, are little affected by the compositional variations studied. This is less common, but not unique, for borate glasses: previous studies of related glasses [46, 51-53] have indicated changes in Mössbauer parameters with different Fe₂O₃ contents. However, equivalent findings to ours were also reported by Akamatsu et al. [22] for xFe₂O₃(80 − x)Bi₂O₃.20.B₂O₃ glasses where (18.2 ≤ x ≤ 40.0), which are closely similar in composition to our glasses. Akamatsu et al. [22] also reported Centre Shift values of 0.32 mm s⁻¹, and only Fe³⁺ was present in their glasses, again in close agreement with our results. Other compositionally-relevant Mössbauer results from literature also include lead borate glasses [51-53] which included high-PbO, low-B₂O₃ glasses containing varying amounts of Fe₂O₃. Sekhon and Kamal [52, 53] observed an approximately monotonic decrease in Fe³⁺ Centre Shift with increasing PbO contents of up to 85mol% PbO in x PbO.(1-x) B₂O₃ glasses containing 10 mol% Fe₂O₃. They attributed this to formation of Fe—O—Pb bonds replacing Fe—O—B bonds with increasing PbO content. Burzo and Ardelean [51] also observed a decrease in Fe³⁺ Centre Shift, here with increasing Fe₂O₃ content from 5 to 50 mol% in xFe₂O₃.1-x(PbO.3B₂O₃) glasses. These works do not readily enable direct and unambiguous comparisons with our data, since Sekhon and Kamal [52, 53] considered a fixed Fe₂O₃ content and boron-rich glasses; and Burzo and Ardelean [45] studied boron-rich glasses. Moreover, the iron in some of their glasses occurred in both Fe²⁺ and Fe³⁺ oxidation states, whereas all of the iron in our glasses occurred as Fe³⁺. Nevertheless, the lack of change in Mössbauer parameters with increasing Fe₂O₃ content of our glasses does enable useful inferences on glass structure. As discussed previously, it is suggested that the large majority (over 80%) of the boron in our glasses occupies trigonal coordination. Consequently the trigonal boron does not require nearby modifier cations for charge balance or stabilisation, and therefore it is less likely that Fe − O − B bonds would be required. However, the tetrahedral fraction of the Fe³⁺ would require nearby cations for charge balance and stabilisation and, moreover, it may be expected that the majority of the Al³⁺ present in these glasses would also require charge balance and stabilisation. This suggests that Fe³⁺ would be more likely to form Fe − O − Bi bonds for any charge balance / stabilisation that may be required, and since there is an abundance of Bi³⁺ in all glasses studied, this may explain the high stability and invariance with composition of the Fe³⁺ sites in these glasses. It is also highly likely that Fe³⁺ − O − Fe³⁺ bonds (clustering) will also be present in these glasses due to their high iron contents. Such behaviour has previously been observed for other borate glasses with lower Fe₂O₃ contents than studied
here [54, 55] and recently, Akamatsu et al. [22], who studied glasses with closely similar compositions to ours, showed that magnetic clusters become more significant as Fe₂O₃ content increases. They also used transmission electron microscopy to show that the magnetic clusters were not nanocrystals but were an Fe-rich amorphous phase. This suggests that sub-nanophase separation occurred in their glasses and, owing to the close compositional similarities between our glasses and those of Akamatsu et al. [22], we can surmise that it may have also occurred in the Fe-doped glasses studied here.
5. Conclusions

Glasses with nominal molar composition \(20\text{B}_2\text{O}_3 - (80-x)\text{Bi}_2\text{O}_3 - x\text{Fe}_2\text{O}_3\) (where \(x = 0-40\)) were successfully prepared by melt-quenching. Spectroscopic studies have confirmed the presence of \(\text{BO}_3\) and \(\text{BiO}_6\) structural units in all glasses, with glass structure apparently little affected by iron, which is solely present in the \(\text{Fe}^{3+}\) oxidation state in all glasses, adopting a distribution of coordinations from 4 to 6. Thermal stability, measured by a number of different parameters, was greatly improved by iron additions, with maximum improvements achieved with 10-20 mol\% \(\text{Fe}_2\text{O}_3\) addition. These improvements are accompanied by modest increases in \(T_g\) and \(T_d\). Consequently it is shown that partial replacement of \(\text{Bi}_2\text{O}_3\) by 10 -20 mol\% \(\text{Fe}_2\text{O}_3\) provides a large improvement in thermal stability of \(\text{B}_2\text{O}_3\)-\(\text{Bi}_2\text{O}_3\) glasses, benefitting the design of new bismuth borate based glasses for low-temperature applications such as sealing.

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