

On the structure of Ge-As-Te-Cu glasses

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Abstract

Short range order in glassy 0.9(Ge_{0.1}As_{0.15}Te_{0.75})-0.1Cu (GATC1) and 0.9(Ge_{0.05}As_{0.55}Te_{0.4})-0.1Cu (GATC2) was studied by neutron- and X-ray diffraction as well as EXAFS (extended X-ray absorption fine structure) measurements at the K-edges of all components. The reverse Monte Carlo simulation technique was used to create models consistent with all experimental datasets. It was found that Cu binds predominantly to Te in GATC1 while Cu-As and Cu-Cu bonding is also significant in GATC2. Ge and As atoms have 4 and 3 Ge/As/Te neighbors in both compositions. In GATC1 the formation of ‘extra’ Te-Te bonds can be observed, similarly to GeTe₄-AgI glasses.

1. Introduction

As-Te alloys can be vitrified over a broad composition range (~20-60 at.% As). A detailed study of As-Te glasses by diffraction techniques and EXAFS revealed that they could be considered as random covalent networks, without any type of preferred bonding [1-3]. The Ge-Te system behaves in a completely different way: liquid Ge-Te alloys can be vitrified only in the vicinity of the eutectic composition (15-20 at. % Ge). A reverse Monte Carlo simulation study using EXAFS and diffraction datasets showed that Ge-Ge bonding is not significant in these glasses [4]. Later this finding was confirmed by EXAFS and X-ray photoelectron

spectroscopy [5]. Given the sensitivity of the experimental techniques applied it means that the Ge-Ge coordination number is not higher than ~ 0.3 , thus amorphous Ge-Te alloys around the eutectic composition should be regarded as chemically ordered.

Ge-As-Te (GAT) alloys can be vitrified between $\sim 80\%$ and $\sim 35\%$ Te content. The different behaviour of As and Ge in binary tellurides is also inherited by the ternary glasses. There is a pronounced ordering around Ge atoms, which prefer Te to As and Ge while As-As bonding is significant even in Te-rich compositions (e.g. $\text{Ge}_{10}\text{As}_{15}\text{Te}_{75}$) [6].

Glass formation in the Ge-As-Te-Cu (GATC) system has been reported recently [7]. GATC glasses have excellent transmission in the IR range and possess high conductivity. This favorable combination of properties makes them suitable for detecting charged biomolecules by electrodepositing them and measuring their IR spectra.

The aim of the present study is the determination of the short range order in GATC. Comparison of the structure of GAT and GATC glasses reveals how Cu atoms modify the Ge-As-Te covalent network. This question is especially interesting because glassy tellurides can host modifiers by entirely different mechanisms. For example, in $0.75\text{GeTe}_4\text{-}0.25\text{AgI}$ the total coordination number of Te is close to 3 and the sum of Te-Ge and Te-Te coordination numbers is 2.76 [8]. Therefore, in this case Ag and I induce a strong rearrangement of the Ge-Te covalent network. On the other hand, in Ge-Ga-Te [4] and Te-rich Ge-Cu-Te [9] glasses the sum of Te-Te and Te-Ge coordination numbers remains close to 2 but the total coordination number of Te is significantly higher than 2, showing that modifying atoms form additional bonds with Te but do not rearrange the host network.

Chemical short range order in an alloy can be characterized by the partial pair correlation functions. The number of these functions in case of an n -component system is $n(n+1)/2$. As four-component glasses have ten partial pair correlation functions the detailed description of the short range order of Ge-As-Te-Cu glasses is a rather challenging goal. Close distances (e.g. Ge-Te, As-Te, Cu-Te) can be separated reliably only by combining various experimental techniques. For this reason we have carried out neutron and X-ray diffraction experiments as well as EXAFS measurements at the K absorption edges of all components. Large scale structural models have been created by fitting simultaneously the 6 experimental datasets in the framework of the reverse Monte Carlo simulation technique [10, 11]. Besides fitting experimental data, this method is also capable of incorporating existing physical and chemical information (e.g. density, preferred coordination numbers, chemical ordering) in the models created. The use of various constraints also gives the possibility of

estimating the uncertainty of coordination numbers or assessing the validity of various structural models, in general.

2. Sample preparation

Both glass samples were prepared using high purity elements (99.999% Ge and Cu, 99.99999% As, and 99.9999% Te) introduced in a silica ampoule which was flame sealed under high vacuum. The melt was homogenized for 8h in a rocking furnace at 850°C. After homogenization, the temperature of the furnace was lowered to 600°C to reduce the vapour pressure and after an isothermal heating for 1 hour the ampoule was quenched in water. The ampoule containing the glass was then immediately annealed at 20°C below the glass transition temperature T_g for 3 hour. The resulting glass rod was recovered by breaking the silica ampoule.

3. Experimental

The neutron diffraction experiment was carried out at the 7C2 liquid and amorphous diffractometer of the Laboratoire Léon Brillouin (Saclay, France). Powdered samples were filled into thin walled vanadium sample holders (diameter: 6 mm, wall thickness: 0.1 mm). 0.9(Ge_{0.1}As_{0.15}Te_{0.75})-0.1Cu was measured with the old ‘banana’ detector filled with BF₃. The new 3He-filled detector system was used to measure 0.9(Ge_{0.05}As_{0.55}Te_{0.4})-0.1Cu. Raw data were corrected for background scattering and detector efficiency. The offset of the detector position and the wavelength of the incident neutron beam (0.72 Å) were determined by measuring a Ni standard.

The X-ray diffraction and EXAFS experiments are described in ref. [6].

4. Reverse Monte Carlo simulation

The reverse Monte Carlo simulation technique was used to generate particle configurations fitting all experimental datasets and using available physical information. Simulation boxes contained 20000 atoms.

Density measurements of Ge₁₅Te₈₅-Cu glasses [12] revealed that the partial molar volume of the Ge₁₅Te₈₅ host matrix (i.e. the volume in which 0.15 mole Ge and 0.85 mole Te can be found) does not depend significantly on the Cu content (up to 10 atomic %). Molar volume of GATC1 was therefore estimated by multiplying the molar volume of Ge₁₀As₁₅Te₇₅

[6] by 0.9. Similar measurements showed that the partial molar volume of $\text{As}_{50}\text{Te}_{50}$ is $\sim 1.8\%$ greater in $\text{As}_{50}\text{Te}_{50}\text{-}10\%\text{Cu}$ than in the undoped binary glass [12]. It is reasonable to assume that the ‘breathing’ of the host matrix increases with decreasing Te content due to the decreasing average distance of the atoms. Thus the molar volume of GATC2 was taken to be only 6% smaller than that of $\text{Ge}_{50}\text{As}_{50}\text{Te}_{40}$ (see Table 1). Test runs revealed that small ($\pm 3\%$) changes of the density have no significant effect on short range order parameters.

Minimum interatomic distances are listed in Table 2. Backscattering factors needed to calculate the model EXAFS curves from partial pair correlation functions [13] were obtained by the feff8.4 programme [14]. Raw $\chi(k)$ data were filtered in two steps: first $k^3\chi(k)$ was forward Fourier-transformed into r-space using a Kaiser-Bessel window ($\alpha=1.5$). The k-range of transformation was around $1.8 \text{ \AA}^{-1}$ - $14 \text{ \AA}^{-1}$ for the As and Te K edges and around $1.8 \text{ \AA}^{-1}$ - $12.5 \text{ \AA}^{-1}$ for the Ge and Cu edges. The resulting r-space data were backtransformed using a rectangular window (usually over the r-space range $1.2 \text{ \AA}$ - $3.2 \text{ \AA}$).

5. Results and discussion

5.1. $0.9(\text{Ge}_{0.1}\text{As}_{0.15}\text{Te}_{0.75})\text{-}0.1\text{Cu}$

First the environment of Ge atoms was investigated. It was found that the elimination of Ge-Ge, Ge-As and Ge-Cu bonds does not influence the quality of fits. Therefore Ge atoms were allowed to have only Te neighbors and the Ge-Te coordination number was constrained to be 4. The Ge-Te bond length ($2.60 \pm 0.02 \text{ \AA}$) is in a good agreement with literature values [4, 6, 15, 16].

Arsenic binds mostly to Te but As-As bonding also improved fit qualities. The total coordination number of As was constrained to be 3. N_{AsAs} , the As-As coordination number is 0.33 ± 0.2 which is very close to the corresponding value (0.39) of $\text{Ge}_{10}\text{As}_{15}\text{Te}_{75}$ glass [6].

Cu has on the average 3.37 ± 0.3 Te neighbors. The mean Cu-Te distance is $2.56 \pm 0.02 \text{ \AA}$, which agrees with the value found in amorphous GeCu_2Te_3 [17]. It was established that the fit of the Cu K-edge EXAFS data was improved upon allowing Cu-Cu bonding. However, the position of the first peak of $g_{\text{CuCu}}(r)$, the Cu-Cu partial pair correlation function, was strongly sensitive to the parameters of the fit (e.g. fitting range, weight of the Cu EXAFS dataset). Therefore it has been concluded that Cu-Cu bonding in the model is an artefact and the real Cu-Cu coordination number is most likely below the sensitivity of our approach.

The total coordination number of Te is 2.74 ± 0.2 . The average number of Cu atoms around Te is 0.50 ± 0.1 while the number of ‘covalent’ neighbours (Ge, As, Te) is 2.24 ± 0.2 .

Comparison with $\text{Ge}_{10}\text{As}_{15}\text{Te}_{75}$ reveals that the first minimum of $g_{\text{TeTe}}(r)$ becomes less well defined upon adding Cu (Fig. 3). The shoulder at 2.9 Å suggests that ‘extra’ Te atoms appear in the first coordination sphere of Te and increase the average coordination number of Te by forming a third covalent bond, similarly to the $\text{GeTe}_4\text{-AgI}$ system [8].

5.2. $0.9(\text{Ge}_{0.05}\text{As}_{0.55}\text{Te}_{0.4})\text{-}0.1\text{Cu}$

As this composition is strongly Te-poor Ge cannot be fourfold coordinated without significant Ge-As bonding (see Table 2). Nevertheless, Ge-Te bonds are clearly preferred to Ge-As ones. Neither Ge-Ge nor Ge-Cu bonding was found to improve fit quality therefore these pairs were forbidden by raising the corresponding minimum interatomic distances (Table 2). Besides Ge, arsenic is mostly surrounded by Te and As. However, test runs showed that Cu also binds to arsenic. The Cu-As coordination number is 1.43 ± 0.7 . Given the high percentage of As, this value still indicates that Cu-Te bonds are strongly preferred to Cu-As ones. The Cu-Cu coordination number is 1.83 ± 1 . The individual uncertainties of Cu-X coordination numbers are rather high but they largely cancel each other, similarly to the case of GATC1. Therefore the total coordination number of Cu is relatively well defined (4.45 ± 0.4). Though Te-Te bonding was allowed, the Te-Te coordination number is only 0.03. This value is well below the sensitivity of our approach therefore Te-Te bonding is not significant in GATC2.

5.3. Comparison with the structure of other amorphous chalcogenides

The Ge-Te distance is 2.60 ± 0.02 Å both in GATC1 and GATC2. The same bond length was found in several Ge-Te based glasses [4, 6, 15, 16] showing that the strength of Ge-Te bonds is not strongly sensitive to the composition. This is definitely not true for the Te-Te distance that shows a great variation even in amorphous Ge-Te films, with a well-defined maximum around $\text{Ge}_{20}\text{Te}_{80}$ [18]. The Te-Te distance also seems to depend on how the modifier binds to the atoms of the host network: in case of covalent bonding (e.g. Se and I) the bond length becomes smaller (2.70-2.73 Å) while Ga increases the mean Te-Te distance to 2.79-2.80 Å [4]. In GATC1 and in $\text{Ge}_{10}\text{As}_{15}\text{Te}_{75}$ the Te-Te bond length is 2.78 ± 0.02 Å and 2.76 ± 0.02 Å, respectively.

Cu-Te distances also lie in a rather narrow range (2.56-2.58 Å) in GATC1, GATC2, and GeCu_2Te_3 [17]. The electronegativity of Cu is 1.9 while that of Te is 2.1. These are very close, hence the Cu-Te bond is expected to be covalent. This is confirmed by the fact that the investigated Ge-As-Te-Cu glasses are amorphous semiconductors with a bandgap around 1 eV [7], which is much smaller than the typical values of ionic compounds.

On the other hand, the scattering of Cu-Cu bond lengths is rather large (2.55-2.81 Å). We believe that this is only partly due to the somewhat higher uncertainty of Cu-related structural parameters. The Cu-Cu distance is strongly sensitive to the environment of Cu atoms. For example in Cu₂Te it is as short as 2.446 Å [19] while in crystalline Cu_{1.4}Te (rickardite) nearest Cu-Cu distances are 2.58 Å and 2.81 Å [20]. Cu-Cu distances found in various amorphous and crystalline copper tellurides are listed in Table 5.

6. Conclusions

The structure of 0.9(Ge₁₀As₁₅Te₇₅)-0.1Cu and 0.9(Ge₅As₅₅Te₄₀)-0.1Cu chalcogenide glasses has been investigated by diffraction and EXAFS measurements. Experimental data (six datasets for each composition) have been fitted simultaneously by the reverse Monte Carlo simulation technique. Ge is fourfold coordinated in both compositions. In Ge₅As₅₅Te₄₀-10Cu the total coordination number of As is higher than 3 due to the formation of Cu-As bonds. Cu-Cu bonding is significant in Ge₅As₅₅Te₄₀-10%Cu thus the prevalence of CuTe₄ structural units can be ruled out in this composition.

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Table 1. Estimated molar volumes (in cm³) of GATC1 and GATC2 used in the reverse Monte Carlo simulation. The values of the corresponding undoped glasses [6] are also given for comparison.

GATC1	18.32
Ge ₁₀ As ₁₅ Te ₇₅	20.35
GATC2	16.88
Ge ₅ As ₅₅ Te ₄₀	17.96

Table 2. Minimum interatomic distances (in Å) used in the reverse Monte Carlo modelling of 0.9(Ge_{0.10}As_{0.15}Te_{0.75})-0.1Cu and 0.9(Ge_{0.05}As_{0.55}Te_{0.4})-0.1Cu (values in parentheses)

Ge-Ge	Ge-As	Ge-Cu	Ge-Te	As-As	As-Cu	As-Te	Cu-Cu	Cu-Te	Te-Te
3.6	3.6 (2.35)	3.1	2.35	2.35	3.1 (2.65)	2.35	3.1 (2.45)	2.35	2.5

Table 3. Coordination numbers of 0.9(Ge_{0.10}As_{0.15}Te_{0.75})-0.1Cu and 0.9(Ge_{0.05}As_{0.55}Te_{0.4})-0.1Cu obtained by simultaneous fitting of diffraction and EXAFS datasets. Values in bold were obtained by coordination number constraints, therefore no uncertainty is given.

	GATC1	GATC2
N_{GeAs}	-	1.80±0.6
N_{GeTe}	4.00	2.20±0.6
N_{AsGe}	-	0.16±0.05
N_{AsAs}	0.37±0.2	1.48±0.3
N_{AsCu}	-	0.28±0.1
N_{AsTe}	2.62±0.2	1.36±0.3
N_{CuAs}	-	1.40±0.4
N_{CuCu}	-	1.91±0.4
N_{CuTe}	3.37±0.3	1.23±0.4
N_{TeGe}	0.53	0.28±0.08
N_{TeAs}	0.52±0.04	1.87±0.4
N_{TeCu}	0.50±0.1	0.35±0.1
N_{TeTe}	1.19±0.2	0.03
N_{Ge}	4.00	4.00
N_{As}	2.99	3.28±0.1
N_{Cu}	3.37±0.3	4.54±0.3
N_{Te}	2.74±0.2	2.53±0.2

Table 4. Nearest neighbour distances (in Å) found in 0.9(Ge_{0.10}As_{0.15}Te_{0.75})-0.1Cu and 0.9(Ge_{0.05}As_{0.55}Te_{0.4})-0.1Cu and in some Ge-Te, Ge-As-Te and Ge-Cu-Te glasses

	GATC1	GATC2	Ge ₂₀ Te ₈₀ [18]	GAT1[6]	GAT2[6]	GeCu ₂ Te ₃ [17]
r_{GeAs}	-	2.47±0.02	-	-	-	
r_{GeTe}	2.60±0.02	2.60±0.02	2.60±0.02	2.60±0.02	2.61±0.02	2.61±0.02
r_{AsAs}	2.45±0.02	2.45±0.02	-	2.45±0.02	2.43±0.02	

r_{AsCu}	-	2.76 ± 0.02	-	-	-	
r_{AsTe}	2.60 ± 0.02	2.62 ± 0.02	-	2.60 ± 0.02	2.62 ± 0.02	
r_{CuCu}	-	2.62 ± 0.04	-	-	-	2.55 ± 0.02
r_{CuTe}	2.56 ± 0.02	2.57 ± 0.02	-	-	-	2.58 ± 0.02
r_{TeTe}	2.78 ± 0.02	-	2.76 ± 0.02	2.76 ± 0.02	2.80 ± 0.03	2.75 ± 0.02

Table 5. Cu-Cu distances in some crystalline and amorphous copper tellurides (in Å)

Composition	r_{CuCu}
Cu_2Te [19]	2.446
$\text{Cu}_{1.4}\text{Te}$ (rickardite) [20]	2.58, 2.81
$\text{Cu}_{1.79}\text{Te}$ (weissite) [21]	2.493-2.722
BaCu_2Te_2 [22]	2.847
TlCu_3Te_2 [23]	2.474-2.890
GeCu_2Te_3 [17]	2.58 ± 0.03
$\text{Ge}_5\text{As}_{55}\text{Te}_{40}$ -10%Cu	2.62 ± 0.04

Figure captions

Figure 1. Comparison of measured diffraction and filtered experimental EXAFS datasets of $0.9(\text{Ge}_5\text{As}_{55}\text{Te}_{40})\text{-}0.1\text{Cu}$ (symbols) and the model curves (solid lines) obtained by reverse Monte Carlo simulation.

Figure 2. Partial pair correlation functions of $0.9(\text{Ge}_{10}\text{As}_{15}\text{Te}_{75})\text{-}0.1\text{Cu}$ (solid lines) and $0.9(\text{Ge}_5\text{As}_{55}\text{Te}_{40})\text{-}0.1\text{Cu}$ (symbols).

Figure 3. Comparison of the Te-Te partial pair correlation functions of $\text{Ge}_{10}\text{As}_{15}\text{Te}_{75}$ [6] and $0.9(\text{Ge}_{10}\text{As}_{15}\text{Te}_{75})\text{-}0.1\text{Cu}$.

Figure 1.

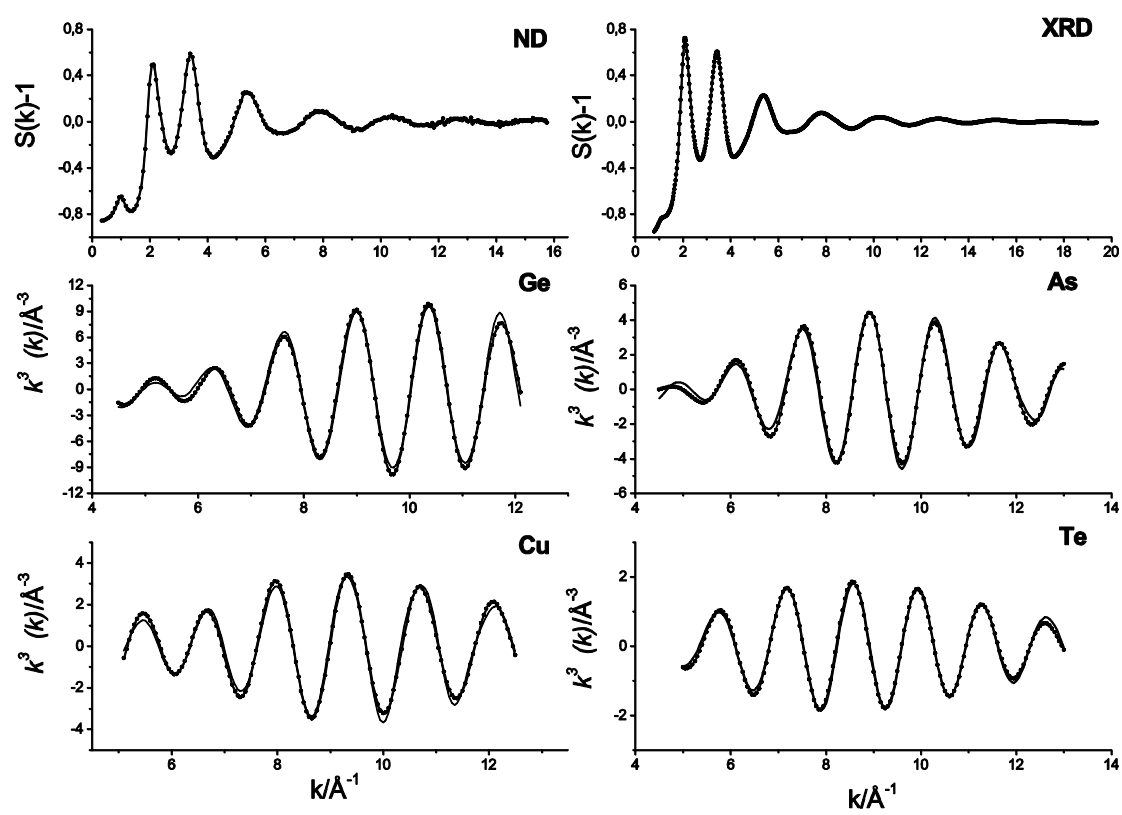


Figure 2.

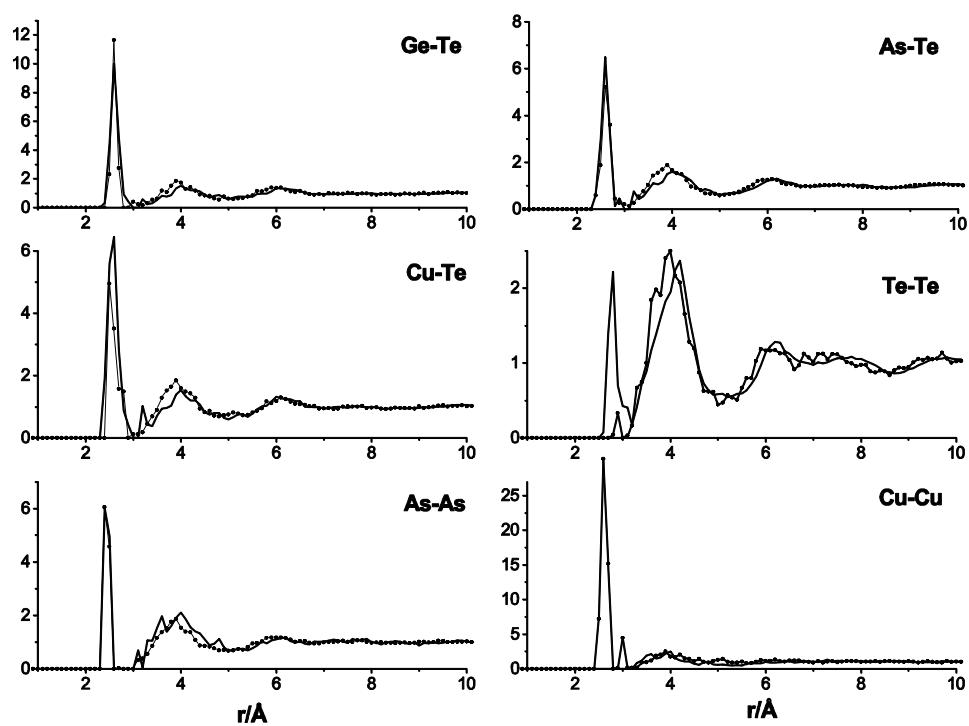


Figure 3.

