Short range order and topology of Ge Ga Te100-2 glasses

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To cite this version:

I. Pethes, A. Piarristeguy, Annie Pradel, S. Michalik, R. Nemausat, et al.. Short range order and topology of Ge Ga Te100-2 glasses. Journal of Alloys and Compounds, Elsevier, 2020, pp.155097. 10.1016/j.jallcom.2020.155097. hal-02549306

HAL Id: hal-02549306
https://hal.archives-ouvertes.fr/hal-02549306
Submitted on 9 Dec 2020

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Abstract: Chemical short range order and topology of GexGaxTe100-2x glasses was investigated by neutron- and x-ray diffraction as well as Ge and Ga K-edge extended x-ray absorption fine structure (EXAFS) measurements. Large scale structural models were obtained by fitting experimental datasets simultaneously with the reverse Monte Carlo simulation technique. Unconstrained models (relying only on experimental data and basic physical information) give 3.9 - 4.1 for the average coordination number of Ge atoms, while the average number of neighbors of Ga atoms scatters around 3.8. The average coordination number of Te atoms is significantly higher than 2 for x = 12.5 and 14.3. It is found that the vast majority of MTe4 (M=Ge or Ga) tetrahedra have at least one corner sharing MTe4 neighbor.
Dear Professor Schultz,

In this submission package, please find our manuscript

'Short range order and topology of Ge$_{x}$Ga$_{1-x}$Te$_{100-2x}$ glasses'

by

PETHES, Ildikó,
PIARRISTEGUY, Andrea,
PRADEL, Annie,
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We kindly ask you to consider the manuscript for publication in the Journal of Alloys and Compounds.

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Thank you very much for your help and kind assistance.

Sincerely,
Ildikó Pethes
Prime Novelty Statement

1. The structure of Ge$_x$Ga$_{1-x}$Te$_{100-x}$ ($x = 7.5, 10, 12.5, 14.3$) glasses has been investigated by combining neutron- and X-ray diffraction, and EXAFS measurements in the same structural model.
2. Our results reveal that the structure of these glasses is characterized by strong chemical ordering.
3. Ge and Ga atoms have 4 Te nearest neighbors.
4. The average coordination number of Te atoms increases with Ga content and it is significantly higher than 2 for $x = 12.5$ and 14.3.
5. The vast majority of Ge(Ga)Te$_4$ tetrahedra have at least one corner sharing Ge(Ga)Te$_4$ neighbor.

It is original research and neither this work itself nor parts of it have been published before or submitted to any other journal.

Sincerely yours,

Ildikó Pethes

corresponding author

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Institute for Solid State Physics
Short range order and topology of Ge$_x$Ga$_x$Te$_{100-2x}$ glasses

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Abstract
Chemical short range order and topology of Ge$_x$Ga$_x$Te$_{100-2x}$ glasses was investigated by neutron- and x-ray diffraction as well as Ge and Ga K-edge extended x-ray absorption fine structure (EXAFS) measurements. Large scale structural models were obtained by fitting experimental datasets simultaneously with the reverse Monte Carlo simulation technique. Unconstrained models (relying only on experimental data and basic physical information) give 3.9 - 4.1 for the average coordination number of Ge atoms, while the average number of neighbors of Ga atoms scatters around 3.8. The average coordination number of Te atoms is significantly higher than 2 for $x = 12.5$ and 14.3. It is found that the vast majority of MTe$_4$ (M=Ge or Ga) tetrahedra have at least one corner sharing MTe$_4$ neighbor.

Introduction
Due to their broad infrared transmission window glassy tellurides are extensively used in various fields of IR optics. The general strategy to find tellurides with excellent glass forming ability is to add a third component to the prototype Ge-Te system. Alloys with Ge-X-Te (X = Ga, As, Se, I, Ag, AgI) composition often possess a broad supercooled liquid region that

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makes it possible to shape bulk infrared lenses or draw fibers transmitting up to at least 18 μm [1 - 3]. It has been shown recently that in binary \( \text{Ge}_x\text{Te}_{100-x} \) \((14.5 \leq x \leq 23.6)\) glasses the total average coordination numbers of Ge and Te atoms are – within the experimental uncertainty – 4 and 2, respectively [4]. It has also been revealed that Ge-Te glasses are chemically ordered: Ge-Te bonds are clearly preferred to Ge-Ge ones, even if Ge-Ge bonds can be found in \( \text{Ge}_{23.6}\text{Te}_{76.4} \). Alloying affects the structure of the host Ge-Te network in different ways. Se and I bind predominantly to Ge and do not change the average coordination numbers of Ge and Te [5, 6]. In Te-poor compositions As atoms bind to Ge, As and Te atoms but the average coordination numbers of Ge and Te atoms do not change here either [7]. On the other hand, in \( \text{GeTe}_4-\text{AgI} \) glasses the average coordination number of Te atoms is significantly higher than 2 even if only Ge/Te neighbors are taken into account [8]. Therefore, the topology of the host Ge-Te network changes significantly upon adding AgI.

The first experimental study of Ge-Ga-Te glasses combining diffraction techniques and EXAFS in the framework of reverse Monte Carlo (RMC) simulation technique [5] reported that the total coordination number of Te is 2.36 ± 0.15 while the average number of neighbors of Ga atoms is about 3 in \( \text{Ge}_{11.1}\text{Ga}_{11.1}\text{Te}_{77.8} \) (the coordination number of Ge atoms was constrained to be 4). As the average coordination number of Ga was reported to be around 4 in several amorphous systems (e.g. \( \text{Ga}_{50}\text{Se}_{50} \) [9], Ga-doped \( \text{Ge:H} \) [10], \( \text{CsCl–Ga}_2\text{S}_3 \) [11] and \( \text{CsCl–La}_2\text{S}_3–\text{Ga}_2\text{S}_3 \) [12]) the above study was followed by a further investigation of \( \text{Ge}_{11.1}\text{Ga}_{11.1}\text{Te}_{77.8} \).

In the experimentally constrained density functional (DFT) study of Voleská et al [13] the starting configuration of DFT simulation was obtained by fitting diffraction and EXAFS datasets simultaneously with RMC. The DFT-optimized configuration was then ’experimentally refined’ again by RMC by using the DFT bond angle distributions as constraints. This configuration reasonably reproduced the experimental data and had a total energy only 33.8 meV/atom higher than that of the original DFT structure. The average coordination numbers of Ga, Ge and Te atoms were 4.08, 3.77 and 2.59, respectively.

While the coordination number of Ga is rather close to the values found in refs. [9 - 12] the average number of neighbors of Te atoms is significantly higher than the experimentally determined coordination number \(2.36 \pm 0.15\). We note here that due to its high concentration in \( \text{Ge}_{11.1}\text{Ga}_{11.1}\text{Te}_{77.8} \) the average coordination number of Te can be deduced from experimental data with a relatively low uncertainty.

The discrepancy of experimental (RMC) and DFT values is due the shallow minimum of the Te-Te partial pair correlation function in the DFT-generated model. More recent DFT studies
emphasized the importance of the choice of exchange-correlation functionals and the proper treatment of van der Waals interactions [14 - 16]. It was demonstrated that by using Becke-Lee-Yang-Parr (BLYP) exchange-correlation functional and van der Waals forces in modelling amorphous tellurides some problems of earlier DFT simulations (e.g. high number of Ge atoms in octahedral environment in a covalent system, too high bond distances) can be avoided. The total coordination number of Te in GeTe₄ is also closer to 2 though the deviation from the experiment-based value is still significant (2.31 vs. 2.00 ± 0.1 in Ge₁₈.₇Te₈₁.₃ [4]). In case of Ge-Ga-Te glasses the main difficulty of experimental structure determination is that Ga and Ge possess similar scattering properties both for X-ray and neutron diffraction (Z₉e=32, Z₉a=31, b₉e=8.185 fm, b₉a=7.288 fm, where Z is the atomic number and b is the coherent neutron scattering length). As Ge and Ga are neighboring elements, Ga K-edge EXAFS signal is limited by the Ge K absorption edge. Another problem is that the mean Ga-Te nearest neighbor distance is between the Ge-Te and Te-Te bond lengths [14], therefore Ga-Te peak parameters (especially the coordination number) are more sensitive to the ‘cross talk’ between overlapping peaks.

Even if the uncertainty of structural parameters is relatively large for a single composition, reliable information can be obtained from experimental data by measuring a concentration series. For this reason, we studied the structure of GeₓGa₁₋ₓTe glasses by combining X-ray and neutron diffraction data with Ge- and Ga K-edge EXAFS measurements in the framework of the reverse Monte Carlo simulation technique. Short range order parameters of Ge-Ga-Te glasses are compared with those of amorphous Ge-Te, Ge-Ga-S and Ge-Ga-Se alloys as well as with models of Ge-Ga-Te glasses obtained by ab initio molecular dynamics.

**Experimental**

**Sample preparation**

Four Ge-Ga-Te glasses of nominal compositions Ge₇.₅Ga₇.₅Te₈₅, Ge₁₀Ga₁₀Te₈₀, Ge₁₂.₅Ga₁₂.₅Te₇₅, and Ge₁₄.₃Ga₁₄.₃Te₇₁.₄ were used for both neutron and X-ray experiments. Starting elements from high-purity, germanium pellets (99.999%, Goodfellow), gallium ingots (99.9995%, Sigma-Aldrich), and tellurium ingots (99.9999%, Sigma-Aldrich) were first weighed in stoichiometric quantities (for a total batch of ~3 g) and introduced in a cylindrical silica tube (11 mm inner diameter, 1 mm thick). The tube was subsequently evacuated under secondary vacuum (10⁻⁵ mbar), sealed and heated up at 1220 K in a furnace with a low heating rate of 10 K/h. The molten batch was held at this temperature for three days and finally quenched in a salt–ice-water after an annealing step of two days at 1073 K.
Neutron diffraction (ND) measurements were carried out at the 7C2 diffractometer of LLB (Saclay, France). The wavelength of incident neutrons was 0.723 Å. Powdered samples were filled into vanadium sample holders of 6 mm diameter and 0.1 mm wall thickness. The wavelength and detector position were determined by measuring a standard Ni powder sample. Raw data were corrected for background scattering and detector efficiency.

High energy X-ray diffraction (XRD) measurement was carried out at the Joint Engineering, Environmental and Processing (I12-JEEP) beamline at Diamond Light Source Ltd (UK). The size of the monochromatic beam was 0.3 × 0.3 mm². A CeO₂ reference sample (NIST Standard Reference Material 674b) was measured at different distances to determine the energy of the incident beam, the sample-to-detector distance, the position of the beam centre and the tilt of the detector. The wavelength of the incident beam and the sample-to-detector distance were 0.1255 Å (98.768 eV) and 336 mm, respectively. Collected 2D diffraction data were integrated into reciprocal-space using the DAWN software [17]. X-ray structure factor, $S_X(Q)$, were extracted from integrated raw data using the PDFGetX2 software [18].

Ge and Ga K-edge EXAFS spectra were measured in fluorescence mode at beamline P65 of the Petra III source. Samples were finely ground, mixed with cellulose and pressed into tablets. Monochromatic radiation was obtained by a Si(111) double crystal monochromator. $\chi(k)$ curves were obtained using the Viper program [19]. Raw $\chi(k)$ signals were first forward Fourier-transformed using a Kaiser-Bessel window. The resulting $r$-space curves were back transformed using a rectangular window over 1.1-2.4 Å.

**Reverse Monte Carlo simulations**

The reverse Monte Carlo (RMC) method [20] is robust tool to obtain large three-dimensional structural models consistent with the supplied (experimental and/or theoretical) data sets. It can be used with any quantity that can be obtained from the atomic coordinates, such as total structure factors from ND or XRD experiments or EXAFS curves. A strength of the method is that the data sets can be fitted simultaneously. During the simulation particles are moved around randomly to minimize the differences between experimental and model curves. Finally particle configurations compatible with all fitted data sets (within the experimental error) are obtained. From these configurations short range order parameters (partial pair correlation functions, average coordination numbers etc.) can be calculated.
The RMC++ code [21] was used to produce structural models by fitting simultaneously the experimental data sets. The EXAFS backscattering coefficients were calculated by the FEFF8.4 program [22].

The investigated samples, their estimated densities and the fitted data sets are collected in Table 1. Densities were estimated using literature values of amorphous GeₓTe₁₀₀₋ₓ [23, 24] and GaₓGeₓTe₁₀₀₋ₓ₋ᵧ glasses [25 - 29]. The simulation boxes contained 10000 atoms for the test runs and 40000 atoms for the final results. Initial configurations were obtained by placing the atoms randomly in the boxes and moving them around to satisfy the minimum interatomic distance (cutoff) requirements. Starting values of the cutoff distances were usually around 85-90% of the sum of the corresponding atomic radii (\(r_{\text{Ge}} \approx 1.25 \text{ Å}, \ r_{\text{Ga}} \approx 1.3 \text{ Å}, \ r_{\text{Te}} \approx 1.4 \text{ Å}\)) [30], the final values are collected in Table 2. Ge-Te, Ga-Te and Te-Te bonds were allowed in all simulation runs. In the investigated samples the amount of Te atoms is more than twice of the amount of Ge + Ga atoms, thus besides the formation of MTe₄ (or MTe₃) units (M=Ge or Ga) Te-Te pairs must also be present, even if heteronuclear Ge-Te and Ga-Te bonds are preferred. The necessity of M-M type bonds was investigated by test runs for the highest M content \(\text{Ge}_{14.3}\text{Ga}_{14.3}\text{Te}_{71.4}\) sample (see below). Allowing them had either no effect on fit quality or resulted in M-M type \(g(r)\) functions with an artificially split first peak having maxima around 2.5 Å and 2.8 Å. From these test runs it was concluded that M-M type coordination numbers in glassy \(\text{Ge}_{14.3}\text{Ga}_{14.3}\text{Te}_{71.4}\) are around or below the sensitivity of our method (about 0.3-0.4) and even smaller in the other compositions investigated.

In the final models all M-M type bonds were forbidden by using cutoff values higher than the expected bond lengths. In all simulation runs some low coordination numbers of the atoms (0 for Te, 0 and 1 for Ga, and 0, 1 and 2 for Ge) were eliminated. In the so called 'unconstrained' models only the above coordination constraints were used. The quality of the fits of different models were compared via their 'goodness-of-fit' (\(R\)-factor) values:

\[
R = \left( \frac{\sum_i (s_{\text{mod}}(Q_i) - s_{\text{exp}}(Q_i))^2}{\sum_i s_{\text{exp}}^2(Q_i)} \right)^{\frac{1}{2}}
\]

(1)

Here \(Q_i\) are the experimental points while 'mod' and 'exp' refer to model and experiment, respectively. Similar expression is valid for the EXAFS curves.

**Results and discussion**
The experimental total structure factors \((S(Q))\) and filtered, \(k^3\)-weighted EXAFS curves \((k^3\chi(k))\) are shown in Figures 1-4. Also shown are the fits of the unconstrained final models, in which only the Ge-Te, Ga-Te and Te-Te bonds were allowed Figs. 1-4.

Partial pair correlation functions \((g_\rho(r))\) obtained for the unconstrained model are shown in Fig. 5 while bond lengths and average coordination numbers are collected in Tables 3 and 4.

**Nearest neighbor distances**

The Ge-Te bond distances are around 2.60 Å, while the Ga-Te bond length is 2.62-2.63 Å for all glasses except the Ge\(_{10}\)Ga\(_{10}\)Te\(_{80}\) sample for which no EXAFS data were available. For this composition longer Ge-Te and Ga-Te distances are (can be) compensated by a shorter Te-Te distance length showing that diffraction data without EXAFS cannot completely separate Ge-Te, Ga-Te and Te-Te distances. For the other compositions the \(r_{\text{GeTe}}\) value agrees well with previous results: in amorphous Ge\(_x\)Te\(_{100-x}\) \(r_{\text{GeTe}} = 2.59\) Å was found by ND [31], 2.59-2.62 Å by EXAFS [32 - 36], 2.58-2.61 Å by combining diffraction, EXAFS and RMC techniques [4, 8, 24, 37, 38], 2.6 Å by anomalous X-ray scattering and RMC [39]. 2.60-2.64 Å Ge-Te bond distances were found in amorphous Ge-Sb-Te [40 - 42], 2.60 Å in Ge-As-Te [7] and 2.60-2.63 Å in Ge-Ga-Te glasses [5, 26].

Early density functional molecular dynamics simulations (DFT) resulted in longer Ge-Te bond lengths: 2.70 – 2.78 Å in Ge\(_x\)Te\(_{100-x}\) and Ge\(_x\)Sb\(_y\)Te\(_{100-x-y}\) systems [23, 43 - 45]. The combination of DFT simulations with RMC refinement gave results closer to the experimental values: 2.65 Å in Ge-Ga-Te [13], 2.58 Å in Ge\(_{15}\)Te\(_{85}\) [46]. Recent DFT simulations using the Becke-Lee-Yang-Parr (BLYP) exchange-correlation functional instead of Perdew-Burke-Ernzerhof (PBE), and especially with van der Waals dispersion forces included, have shown improved agreement with experimental values: 2.64-2.66 Å with PBE in Ge\(_x\)Te\(_{100-x}\) [15, 47 - 49], 2.59-2.62 Å with BLYP in Ge\(_x\)Te\(_{100-x}\) [15], 2.63 Å in Ge\(_2\)Sb\(_2\)Te\(_5\) [50] and 2.59 Å in Ge\(_{15}\)Ga\(_{10}\)Te\(_{75}\) [14].

The \(r_{\text{GaTe}}\) value (2.62-2.63 Å) is the same as it was found experimentally in Ge-Ga-Te glasses by EXAFS [26] and by combining experimental (diffraction, EXAFS) data with simulation (RMC and density functional) [13]. Somewhat longer bond length was found in Ref. [14] (2.67 Å) with first principles molecular dynamics simulation (FPMD).

The Te-Te bond length is around 2.77 Å, except again for the Ge\(_{10}\)Ga\(_{10}\)Te\(_{80}\) sample. Te-Te distances reported in the literature have a broad distribution: in amorphous Ge\(_x\)Te\(_{100-x}\) 2.76 Å was measured by ND [31], 2.77-2.82 Å by EXAFS [33, 34], 2.70-2.79 Å by combination of diffraction, EXAFS and RMC [4, 5, 7, 8, 24, 37, 38], 2.73 Å by anomalous X-ray scattering and RMC [39]. Te-Te distances around 2.79-2.80 Å were reported in Ge-Ga-Te [5], 2.77 Å in
Ge-As-Te [7], 2.77-2.79 Å in Ge-Te-Ag-I glasses [8] by combination of diffraction, EXAFS and RMC.

First principles molecular dynamics simulations result in longer bond lengths: 2.87 Å [43], 2.85 Å [45], 2.90 Å [47, 48], 2.89 Å [49]. Shorter distances were obtained by DFT simulations with RMC refinement: 2.83 Å [13], 2.74 Å [46]. Treatment of van der Waals forces (VdW) seems to be important here as well: 2.83 -2.84 Å was obtained with and 2.88-2.89 Å without VdW in GeTe₄ [15] and 2.81 Å in Ge₁₅Ga₁₀Te₇₅ with BLYP+VdW [14].

**Average coordination numbers**

The average coordination number of Ge is around 4, as it is expected (and was reported earlier by experiments and simulations as well [4, 5, 7, 8, 13-15, 23, 26, 31, 33, 37, 39, 41-43, 45-51]).

The average coordination number of Ga atoms is also close to 4, as was found in several Ge-Ga-Ch (Ch=S, Se, Te) glasses: experimentally in Ge-Ga-S glasses [52 - 59], in Ge-Ga-Se glasses [54, 60 - 65], in amorphous Ge-Ga-Te [13, 26]. Recent FPMD simulations on 80GeSe₂-20Ga₂Se₃ glass [66], Ge-Ga-Te liquids [16] and Ge₁₅Ga₁₀Te₇₅ glass [14] also reported Ga coordination numbers around 4.

Test simulation runs were made in which coordination constraints were used to force Ge and Ga atoms to have exactly 4 neighbors (about 95% of the atoms satisfied this requirement). It was found that the quality of the fits of these models was as good as that of the unconstrained model.

The total coordination number of Te increases with increasing Ge/Ga content (see Table 4). It is around 2 for the Ge₇.₅Ga₇.₅Te₈₅ glass and significantly higher than 2 for Ge₁₂.₅Ga₁₂.₅Te₇₅ and Ge₁₄.₃Ga₁₄.₃Te₇₁.₄ (around 2.19 and 2.35, respectively). Test runs were carried out in which the Ge-Te and Ga-Te coordination numbers were constrained to remain 4 and the Te-Te coordination number was forced to decrease so that the total coordination number of Te be equal to 2. The R-factors of the fits for these models were significantly higher (mostly the R-factors of the ND and XRD data sets, with 20-50%), especially for Ge₁₂.₅Ga₁₂.₅Te₇₅ and Ge₁₄.₃Ga₁₄.₃Te₇₁.₄. Besides the deterioration of the fits, the resulting Te-Te partial pair correlation functions exhibit artificially sharp peaks at around 3.1 Å, next to the upper limit of the coordination constraints.

Te coordination number around 2 was found in GeₓTe₁₀₀-x glasses by ND [31] and by EXAFS [34]. Nₓ = 2 was obtained by combination of diffraction and EXAFS experiments with RMC simulations in amorphous GeₓTe₁₀₀-x [5, 24, 7, 8, 4], in Ge-As-Te glasses [7] and in amorphous Ge-Sb-Te [41, 42].
FPMD simulations often result in Te coordination number higher than 2 (e.g. in Ge₄Te₁₀₀₋ₓ [43, 23, 47, 48, 15, 49], in Ge-Sb-Te [43, 51, 45]), the value highly depends on the used exchange correlation functional and on the applying of the VdW forces (see e.g. Ref. [15]: 3.78 PBE, 3.51 PBE+VdW, 2.57 BLYP, 2.31 BLYP+VdW). Recent FPMD simulations proposed Te coordination number 2.9-4.8 in Ge-Ga-Te liquids [16], and 2.19 in Ge-Ga-Te glass [14]. The first minimum of $g_{\text{TeTe}}(r)$ obtained by these simulations is far from zero, thus the second coordination sphere may also contribute to the coordination number of Te.

As in Ge-Te glasses the Te coordination number follows from fitting simultaneously diffraction and EXAFS datasets with RMC simulation [4], the higher coordination of Te in Ge-Ga-Te glasses is due the presence of Ga atoms. It is to be noted that chemical ordering is also different in the two systems. While Ge-Ge bonds can be observed in melt quenched Ge₂₃.₆Te₇₆.₄ [4] no M-M type bonds were found in Ge-Ga-Te glasses. In principle, the investigated Ge-Te and Ge-Ga-Te glasses are all Te-rich therefore M-M bonding could be avoided even if all Te atoms remain twofold coordinated. Still, M-M bonds exist in Ge₂₃.₆Te₇₆.₄ and the coordination number of Te is higher than 2 in Ge-Ga-Te glasses. These two tendencies share a common consequence: they increase the Te-Te coordination number. Further experimental and theoretical studies are needed to see whether this is just a coincidence or a certain number of Te-Te bonds is required by the glassy state due to energetic or kinetic reasons. It is to be noted that $N_{\text{Se}}$ higher than 2 was found in Ge-Ga-Se glasses also by using various techniques (e.g. ND, EXAFS and RMC [64] or anomalous X-ray scattering and RMC [65]). Moreover, $N_{\text{S}}$ coordination number higher than 2 was proposed in amorphous Ge-Ga-S alloys by Raman [57] as well.

Second neighbors

It was found in Ge-Te glasses [4] that GeTe₄ units connect to each other by sharing a corner or an edge (two Ge atoms sharing one or two common Te neighbors). The presence of corner (CS) or edge sharing (ES) MTe₄ tetrahedra was investigated in Ge-Ga-Te samples as well. The analyzed configurations were obtained by constrained simulation runs, in which the Ge and Ga atoms were forced to have 4 Te neighbors. (Images of the configurations of Ge₇.₅Ga₇.₅Te₈₅ and Ge₁₄.₃Ga₁₄.₃Te₇₁.₄ glasses are shown in Figs. 6 and 7.)

The decomposition of the first peak of the M-M partials to contributions of CS and ES tetrahedra and topologically distant pairs is presented for the Ge-Ge pairs of the Ge₁₄.₃Ga₁₄.₃Te₇₁.₄ sample in Fig. 8 (Ga-Ga and Ge-Ga pairs show a similar behavior). It was found that even for the Ge₇.₅Ga₇.₅Te₈₅ sample about 84% of GeTe₄ and GaTe₄ tetrahedra have at least one corner sharing MTe₄ neighbor (see Table 5). This value seems to
be rather high in view of the low Ge/Ga-content of this glass. The average number of Ge and Ga atoms around Te is less than 1 meaning that formation of CS or ES pairs could be avoided, in principle. With increasing M content the number of the CS or ES sharing tetrahedra also increases. For the Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ glass 99.7% of the MTe$_4$ tetrahedra have at least one CS or ES pair. The number of M atoms participating in ES units is around 12% for Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ and 33% for Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$.

**Prepeak in the ND total structure factor**

The neutron diffraction structure factor of Ge$_x$Ga$_x$Te$_{100-2x}$ glasses has a first sharp diffraction peak (FSDP) or prepeak at $q_{\text{max}} \approx 1$ Å$^{-1}$. (A less pronounced peak can be observed in X-ray diffraction structure factor as well.) Peak positions and heights are given in Table 6. The height of the prepeak is defined as $S(q_{\text{max}}) - S(q_{\text{min}})$ where $q_{\text{min}}$ is the first minimum after the prepeak. For Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ there is only a shoulder therefore we used the $q_{\text{min}}$ value of Ge$_{10}$Ga$_{10}$Te$_{80}$. It can be observed that in case of Ge$_x$Ga$_x$Te$_{100-4}$ glasses the height of the prepeak increases with increasing Ge/Ga content.

The connection of medium range order and prepeak intensity is confirmed by comparing the models of Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ obtained with and without fitting neutron diffraction data. Some results of these runs are shown in Fig. 9. It can be observed that models obtained without fitting neutron diffraction data fail to reproduce the prepeak of neutron diffraction structure factors. The other effect of omitting neutron diffraction data from the models is the rather flat first peak of the Ge-Ge partial pair correlation function. A similar behavior can be observed in Ge$_{18.7}$Te$_{81.3}$[4], also shown in Fig. 9. The first peaks of Ge-Ga and Ga-Ga partial pair correlation functions are affected in the same way (not shown). These observations strongly suggest that prepeak in Ge$_x$Ga$_x$Te$_{100-2x}$ and Ge-Te glasses is connected to the well-defined M-M correlations (M = Ge, Ga) manifested in sharp real space M-M peaks. The latter, on the other hand, are the consequence of corner or edge sharing MTe$_4$ tetrahedra [4].

**Conclusions**

Short range order and topology of Ge$_x$Ga$_x$Te$_{100-2x}$ ($x = 7.5, 10, 12.5, 14.3$) glasses was investigated by diffraction techniques and EXAFS. Structural models were obtained by fitting experimental datasets simultaneously in the framework of the reverse Monte Carlo simulation technique. It was shown that Ga and Ge atoms are mostly fourfold coordinated while $N_{\text{Te}}$, the average coordination number of Te increases with Ga content ($N_{\text{Te}} = 2.35 \pm 0.1$ for $x = 14.3$). The majority of Ge/Ga atoms are linked to other Ge/Ga atoms via one or two common Te neighbors forming corner and edge sharing tetrahedra.
Acknowledgment

I. P. and P. J. were supported by NKFIH (National Research, Development and Innovation Office) Grant No. KH 130425. The neutron diffraction experiment was carried out at the ORPHÉE reactor, Laboratoire Léon Brillouin, CEA-Saclay, France. EXAFS measurements were carried out at PETRA III at DESY, a member of the Helmholtz Association (HGF). We thank Diamond Light Source Ltd. for access to beamline I12-JEEP (cm19662-1) that contributed to the results presented here.

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Tables

Table 1 Estimated densities, number densities, fitted experimental data sets.

|        | $\rho$ [g/cm$^3$] | Number density [Å$^{-3}$] | Experimental data sets          |
|--------|-------------------|--------------------------|---------------------------------|
| Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ | 5.6                | 0.0283                   | ND, XRD, Ge, Ga EXAFS           |
| Ge$_{10}$Ga$_{10}$Te$_{80}$  | 5.57               | 0.0288                   | ND, XRD                        |
| Ge$_{12.5}$Ga$_{12.5}$Te$_{75}$ | 5.53               | 0.0293                   | ND, XRD, Ge, Ga EXAFS           |
| Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ | 5.7                | 0.0308                   | ND, XRD, Ge, Ga EXAFS           |

Table 2 Minimum interatomic distances (in Å) used in the reverse Monte Carlo simulation runs

|                  | Ge-Ge | Ge-Ga | Ge-Te | Ga-Ga | Ga-Te | Te-Te |
|------------------|-------|-------|-------|-------|-------|-------|
| Bond allowed     | 2.35  | 2.35  | 2.4   | 2.35  | 2.4   | 2.5   |
| Bond forbidden   | 3.45  | 3.45  | 3.45  |       |       |       |

Table 3 Nearest neighbor distances (in Å). The uncertainty of distances is about ±0.02 Å if EXAFS data sets are also fitted, but it can be higher (about 0.05 Å) in case of the Ge$_{10}$Ga$_{10}$Te$_{80}$, where only the two diffraction measurements were available.

|        | Ge-Te | Ga-Te | Te-Te |
|--------|-------|-------|-------|
| Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ | 2.60  | 2.62  | 2.78  |
| Ge$_{10}$Ga$_{10}$Te$_{80}$  | 2.64  | 2.64  | 2.74  |
| Ge$_{12.5}$Ga$_{12.5}$Te$_{75}$ | 2.60  | 2.63  | 2.77  |
| Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ | 2.61  | 2.62  | 2.77  |

Table 4 Coordination numbers of the investigated glasses obtained by unconstrained simulations.

| Pair (upper limit) | Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ | Ge$_{10}$Ga$_{10}$Te$_{80}$ | Ge$_{12.5}$Ga$_{12.5}$Te$_{75}$ | Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ |
|-------------------|-------------------------------|-------------------------------|---------------------------------|---------------------------------|
| $N_{\text{Ge-Te}}$ (3.1 Å) | 4.0 (-0.4+0.6)               | 4.1 (-0.4+0.7)               | 3.9 (-0.4+0.5)                  | 3.95 (±0.3)                     |
| $N_{\text{Te-Ge}}$ (3.1 Å) | 0.35 (-0.03+0.06)            | 0.51 (-0.05+0.1)             | 0.65 (±0.08)                    | 0.79 (±0.06)                    |
Table 5 Percentage of Ge and Ga atoms participating in corner or edge sharing units.

Uncertainties were determined from 10 simulation runs started from different initial configurations.

|                  | Ge\textsubscript{7.5}Ga\textsubscript{7.5}Te\textsubscript{85} | Ge\textsubscript{10}Ga\textsubscript{10}Te\textsubscript{80} | Ge\textsubscript{12.5}Ga\textsubscript{12.5}Te\textsubscript{75} | Ge\textsubscript{14.3}Ga\textsubscript{14.3}Te\textsubscript{71.4} |
|------------------|---------------------------------------------------------------|---------------------------------------------------------------|---------------------------------------------------------------|---------------------------------------------------------------|
| Neither CS nor ES Ge | 16.5 (±1)                                                   | 6.8                                                           | 1.5                                                           | 0.3 (±0.1)                                                   |
| Neither CS nor ES Ga | 16.5 (±2)                                                   | 5.6                                                           | 1.1                                                           | 0.25 (±0.15)                                                |
| Only CS Ge       | 71.5 (±1.5)                                                   | 75                                                             | 71                                                             | 65 (±1.5)                                                    |
| Only CS Ga       | 71.5 (±2)                                                   | 75                                                             | 71                                                             | 67 (±1)                                                     |
| Only ES Ge       | 3.2 (±0.5)                                                    | 2.9                                                           | 1.6                                                           | 0.8 (±0.2)                                                   |
| Only ES Ga       | 3.2 (±0.5)                                                    | 2.3                                                           | 1.5                                                           | 0.5 (±0.2)                                                   |
| CS and ES Ge     | 9 (±1)                                                       | 15                                                            | 26                                                             | 34 (±2)                                                     |
| CS and ES Ga     | 9 (±1)                                                       | 17                                                            | 26                                                             | 32 (±1)                                                      |

Table 6 Position of the FSDP ($q_{\text{max}}$) and amplitude of the FSDP (see text for definition)

|                  | $q_{\text{max}}$ [Å$^{-1}$] | $q_{\text{min}}$ [Å$^{-1}$] | $S(q_{\text{max}})-S(q_{\text{min}})$ |
|------------------|-----------------------------|-----------------------------|----------------------------------------|
| Ge\textsubscript{7.5}Ga\textsubscript{7.5}Te\textsubscript{85} | 1.01                        | 1.33                        | -0.003                                  |
| Ge\textsubscript{10}Ga\textsubscript{10}Te\textsubscript{80} | 1.01                        | 1.33                        | 0.13                                    |
| Ge\textsubscript{12.5}Ga\textsubscript{12.5}Te\textsubscript{75} | 1.01                        | 1.33                        | 0.198                                   |
| Ge\textsubscript{14.3}Ga\textsubscript{14.3}Te\textsubscript{71.4} | 0.93                        | 1.33                        | 0.256                                   |
Figures

**Figure 1.** ND structure factors (symbols) and fits (lines) of the Ge$_{x}$Ga$_{1-x}$Te$_{100-2x}$ glasses. (The curves are shifted vertically for clarity.)
Figure 2. XRD structure factors (symbols) and fits (lines) of the Ge$_x$Ga$_{100-x}$Te$_{100-2x}$ glasses. (The curves are shifted vertically for clarity.)
Figure 3. $k^3$-weighted, filtered EXAFS spectra at Ge K-edge (symbols) and fits (lines) of the Ge$_x$Ga$_{1-x}$Te$_{100-x}$ glasses. (The curves are shifted vertically for clarity.)
Figure 4. $k^3$-weighted, filtered EXAFS spectra at Ga K-edge (symbols) and fits (lines) of the Ge$_x$Ga$_x$Te$_{100-2x}$ glasses. (The curves are shifted vertically for clarity.)
Figure 5. Partial pair correlation functions of the Ge$_x$Ga$_y$Te$_{100-2x}$ glasses.
Figure 6. A snapshot about a part of the configuration of the Ge$_{7.5}$Ga$_{7.5}$Te$_{85}$ glass obtained by RMC simulation. The Ge, Ga and Te atoms are represented by magenta, blue and grey balls, respectively. Two corner sharing tetrahedra are marked with orange, a chain of Te-Te nearest neighbors is highlighted by red.
**Figure 7.** A snapshot about a part of the configuration of the Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ glass obtained by RMC simulation. The Ge, Ga and Te atoms are represented by magenta, blue and grey balls, respectively. Two edge-sharing GaTe$_4$ tetrahedra are marked by red; chains of corner-shared MTe$_4$ units are highlighted with orange.
Figure 8. Decomposition of $g_{\text{GeGe}} (r)$ of $\text{Ge}_{14.3}\text{Ga}_{14.3}\text{Te}_{71.4}$ glass to contributions from corner (CS) and edge (ES) sharing tetrahedra and topologically distant Ge-Ge pairs.
Figure 9. Comparison of the neutron weighted structure factors (a and c) and the first peak of the Ge-Ge partial pair correlation functions (b and d) of Ge$_{14.3}$Ga$_{14.3}$Te$_{71.4}$ and Ge$_{18.7}$Te$_{81.3}$ glasses obtained from RMC simulations by (red lines) fitting the experimental neutron diffraction data and (blue line) without fitting ND data. Experimental data (symbols) are also shown for reference.
Models of Ge$_x$Ga$_x$Te$_{100-2x}$ glasses consistent with multiple datasets are constructed

The compositions investigated are characterized by strong chemical ordering

Ge and Ga atoms have 4 Te nearest neighbors

Average coordination number of Te atoms increases with Ga content

Majority of the Ge(Ga)Te$_4$ tetrahedra are connected by their corners or edges