

Ultrahigh-pressure polyamorphism in GeO₂ glass with coordination number >6

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Knowledge of pressure-induced structural changes in glasses is important in various scientific fields as well as in engineering and industry. However, polyamorphism in glasses under high pressure remains poorly understood because of experimental challenges. Here we report new experimental findings of ultrahigh-pressure polyamorphism in GeO₂ glass, investigated using a newly developed double-stage large-volume cell. The Ge–O coordination number (CN) is found to remain constant at ~6 between 22.6 and 37.9 GPa. At higher pressures, CN begins to increase rapidly and reaches 7.4 at 91.7 GPa. This transformation begins when the oxygen-packing fraction in GeO₂ glass is close to the maximal dense-packing state (the Kepler conjecture = ~0.74), which provides new insights into structural changes in network-forming glasses and liquids with CN higher than 6 at ultrahigh-pressure conditions.

high pressure | polyamorphism | glass | oxygen packing

Understanding the structural response of network-forming glasses to pressure is of great interest not only in condensed matter physics, geoscience, and materials science, but also in engineering and industry. As prototype network-forming glasses, silica (SiO₂) and germania (GeO₂) have been the most extensively studied (1–5). These two glasses have similar structural change pathways at high pressures. At ambient pressure, both glasses are composed of corner-linked AO₄ tetrahedra, with atom A (Si or Ge) in fourfold coordination (6). Under compression, the coordination gradually changes from 4 to 6 over a wide pressure range [~15–40 GPa for SiO₂ glass (2, 4) and ~5–15 GPa for GeO₂ glass (1, 3, 5)].

A recent study (7) found that evolution of network-forming structural motifs in glasses and liquids at high pressures can be rationalized in terms of oxygen-packing fraction (OPF). Fourfold-coordinated structural motifs in SiO₂ and GeO₂ glasses are stable over a wide range of OPF between 0.40 and ~0.59. The fourfold-coordinated structural motifs become unstable when the OPF approaches the limit of random loose packing of hard spheres (0.55–0.60) (8, 9). When OPF >~0.60, coordination number (CN) gradually increases with OPF to the limit of random close packing (0.64) (8, 9), where CN increases sharply to 6 with almost-constant OPF ~0.64. Higher-pressure data for SiO₂ glass suggest the existence of another stability plateau for sixfold-coordinated structural motifs, with OPF of up to ~0.72 (7).

The highest coordination that has been experimentally determined so far in SiO₂ and GeO₂ glasses is 6. X-ray diffraction measurement for SiO₂ glass confirmed that sixfold-coordination structural motifs are stable up to 100 GPa (4). For GeO₂ glass, X-ray and neutron diffraction data are limited to 18 GPa (1, 3, 5). X-ray absorption spectroscopic measurements were conducted to 64 GPa (10, 11). Ref. 11 showed no major change in X-ray absorption fine structure up to 64 GPa, although a slight discontinuous change in density is observed around 40–45 GPa.

Some simulation studies predicted the existence of structural motifs with CN >6 above ~100 GPa for SiO₂ liquid (12) and glass (13) and above ~60 GPa for GeO₂ glass (13), with no experimental confirmation so far. A study (14) of SiO₂ glass using

Brillouin scattering in a diamond anvil cell (DAC) showed a kink in the pressure dependence of shear-wave velocity at ~140 GPa and was interpreted as evidence of ultrahigh-pressure polyamorphism in SiO₂. However, no structural information is available under such high pressures. In this study, we developed a new double-stage cell, which enables us to study structure of GeO₂ glass at in situ high-pressure conditions up to 91.7 GPa.

Large-volume samples are vital for accurate measurements on the structure of glasses at high pressures using X-ray diffraction because of the weak X-ray scattering from amorphous materials. In such measurements, a large diffraction angle is essential for accurately determining the structure factor with sufficiently large coverage of momentum transfer Q ($Q = 4\pi E \sin\theta / 12.398$, where E is X-ray energy in keV and θ is the diffraction angle), and for high resolution in the reduced pair distribution function in real space. Recently, generation of pressure up to 94 GPa has been achieved in a DAC with 1-mm culet size anvils (15). However, this large-volume DAC is designed for neutron diffraction measurement; it is difficult to apply this apparatus for X-ray diffraction measurement because of limited solid-angle access. Similarly, there have been attempts to generate high pressures by inserting diamond anvils inside multianvil large-volume presses (16, 17). However, multianvil presses generally have even more limited solid-angle access for X-ray diffraction signals.

We have developed a new double-stage Paris–Edinburgh (PE)-type large-volume press to generate high pressures with large sample volume (Fig. 1A). A pair of second-stage diamond anvils is introduced into the first-stage PE anvils. This combination of opposed anvils (both first- and second stage) provides a large opening in the horizontal plane for X-ray diffraction measurement. To reduce absorption of the gasket surrounding the sample, we used a cubic boron nitride + epoxy (10:1 in weight ratio) inner

Significance

A new double-stage large-volume cell was developed to compress large GeO₂ glass samples to near 100 GPa and to conduct multiangle energy-dispersive X-ray diffraction measurement for in situ structure measurements. We find new experimental evidence of ultrahigh-pressure polyamorphism in GeO₂ glass with coordination number (CN) significantly >6. The structural change to CN higher than 6 is closely associated with the change in oxygen-packing fraction. Our results provide direct structural evidence for ultradense network-forming glasses and liquids. The observed ultrahigh-pressure polyamorphism may also exist in other network-forming glasses and liquids as well, such as SiO₂ and other silicate and germanate systems.

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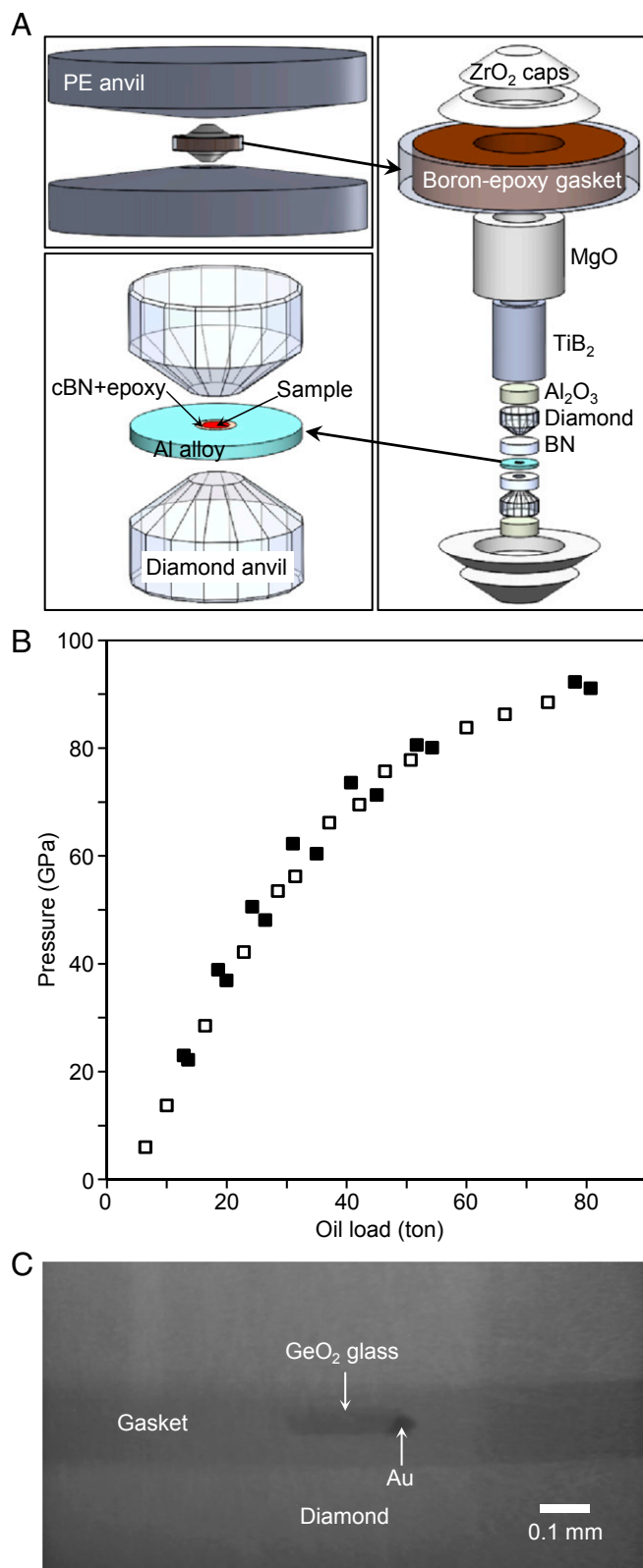


Fig. 1. Newly developed double-stage large-volume cell. (A) Illustration of double-stage large-volume cell design. We use cup-shaped PE-type anvil as the first-stage anvil, and the second-stage diamond anvil is introduced inside the large-volume cell assembly. (B) Pressure generation as a function of oil load of the PE press. Solid symbols represent pressure conditions of structure measurement of GeO_2 glass measured before and after structural measurement. Oil pressure was found to decrease slightly after each measurement, whereas sample pressure increased slightly. Error of pressure is smaller

gasket with an aluminum alloy (7075) outer gasket, both of which are low X-ray absorbing, light-element materials. The diamond anvils had a culet diameter of 0.8 mm. The large culet size allowed us to use large samples 0.3 mm in diameter and 0.15 mm in height. In our study on structure of GeO_2 glass, this new double-stage cell has reached pressures up to 91.7 GPa (Fig. 1B), where sample size was ~ 0.24 mm in diameter and ~ 0.06 mm in height, as determined by in situ X-ray radiography imaging (Fig. 1C).

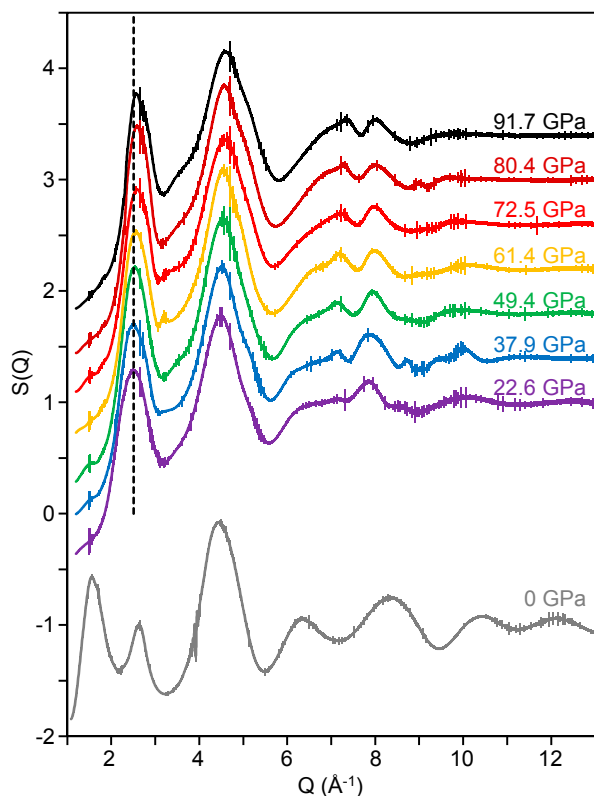
Structure of GeO_2 glass was investigated using the newly designed double-stage PE press with multiangle energy dispersive X-ray diffraction technique at Beamline 16-BM-B, High Pressure Collaborative Access Team (HPCAT) of the Advanced Photon Source (18). Structure factors, $S(Q)$, were obtained up to 13 \AA^{-1} (Fig. 2). The position of the first sharp diffraction peak (FSDP) in $S(Q)$ shows essentially no pressure dependence between 22.6 and 37.9 GPa. Previous studies (3, 19) show that below ~ 10 GPa, FSDP moves rapidly with increasing pressure toward higher Q values; above 10 GPa, the position becomes virtually pressure independent. This is consistent with our observation that the FSDP shifts rapidly from 0 to 22.6 GPa, and then remains almost constant between 22.6 and 37.9 GPa (Fig. 2). Interestingly, with further increase of pressure to 72.5 GPa, the FSDP begins shifting again toward higher Q and then becomes virtually constant once more between 72.5 and 91.7 GPa. Furthermore, a new peak at $Q \sim 7.1\text{--}7.3 \text{ \AA}^{-1}$ begins to emerge at a pressure between 22.6 and 37.9 GPa, with increasing intensity at higher pressures.

Fourier transformation of $S(Q)$ yields real-space pair distribution function (20), $G(r)$ (Fig. 3). The first and second peak of $G(r)$ in GeO_2 glass is generally considered to represent Ge–O and Ge–Ge distance, respectively. Significant changes in peak shape and positions in $G(r)$, particularly in the second peak, are evident. $G(r)$ obtained at 22.6 and 37.9 GPa show a distinct shoulder at the lower “ r ” side of the second peak. This shoulder has been observed in previous studies (1, 3, 5) at pressures higher than ~ 15 GPa. This double-peak feature is considered to represent two Ge–Ge distances in the sixfold-coordinated GeO_2 structure. In contrast, we find that the second peak starts to become a single peak above 49.4 GPa. The second peak width markedly decreases between 37.9 and 49.4 GPa, although the second peak in $G(r)$ at 49.4 and 61.4 GPa still shows a weak shoulder at the low- r side. The second peak becomes a single peak above 72.5 GPa.

Fig. 4 shows the pressure dependence of the first and second peak positions in $G(r)$, with the numerical results summarized in Table 1. We fit the second peak at 22.6–61.4 GPa by two peaks, whereas those at higher pressures are fitted by a single peak. The first peak position in $G(r)$ decreases between 22.6 and 37.9 GPa, whereas it is almost constant or increases slightly between 37.9 and 61.4 GPa. In contrast, above 72.5 GPa, the first peak position decreases markedly to $\sim 1.81 \text{ \AA}$. The second peak position also shows marked change with increasing pressure. The main second peak position at $\sim 3.2 \text{ \AA}$ is almost constant between 22.6 and 37.9 GPa, whereas it decreases markedly with increasing pressure above 49.4 GPa.

Our results reveal that a distinct change in the structure of GeO_2 glass begins at a pressure between 37.9 and 49.4 GPa. Below 37.9 GPa, our structural data are similar to those obtained around 15–18 GPa in previous studies (1, 3, 5). Because the first and second peak positions of $G(r)$ in GeO_2 glass are considered to represent Ge–O and Ge–Ge distances, respectively, we compare the first and second peak positions of $G(r)$ obtained in this

than the size of the symbol. (C) X-ray radiography image of the GeO_2 glass sample and Au pressure standard through gasket at 91.7 GPa.



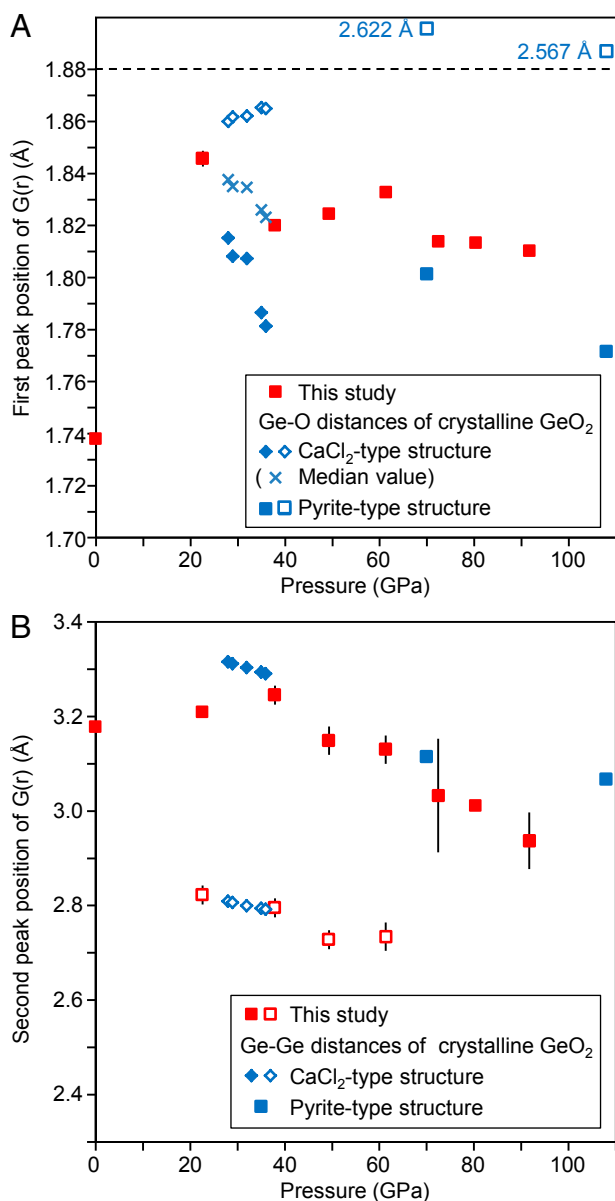


Fig. 4. The first and second peak position of $G(r)$. (A) The first peak position of $G(r)$ of GeO_2 glass obtained in this study (solid red squares), compared with Ge-O bond distance of crystalline GeO_2 with CaCl_2 -type structure (21) (open and solid blue diamonds) and pyrite-type structure (24) (open and solid blue squares). Blue cross symbols represent the median value of the two Ge-O bond distances in CaCl_2 -type GeO_2 . Error is smaller than the size of the symbol. (B) The second peak position of $G(r)$ of GeO_2 glass obtained in this study (red squares), compared with Ge-Ge bond distance of crystalline GeO_2 with CaCl_2 -type structure (21) (open and solid blue diamonds) and pyrite-type structure (24) (solid blue square). Solid red squares represent the main second peak (r2) and open red squares represent shoulder peaks in the second peak at ~ 2.7 – 2.8 Å (r2s) (Table 1). Vertical bars represent errors of the peak positions.

at 91.7 GPa than the extrapolated GeO_2 glass density values. If we adopt the density value of pyrite-type structure crystalline GeO_2 at 72.5–91.7 GPa, CN becomes almost constant around 7. To precisely determine CN of GeO_2 glass and to discuss the change of CN with increasing pressure, precise density data particularly above 70 GPa are required. Nevertheless, our structural results provide strong evidence of GeO_2 glass possessing an ultrahigh-pressure polyamorphic structure with CN > 6 above 49.4 GPa.

A recent study (7) shows that the CN of network-forming structural motifs in oxide glasses and liquids can be rationalized in terms of the OPF. Because of the lack of experimental data with CN > 6, the reported data (7) are limited to $3 < \text{CN} < 6$. We can now extend the relationship between CN and OPF to ultradense glasses and liquids. We calculate OPF (η_O) by using the same method as given by ref. 7 [$\eta_O = V_O/\rho c_O$, $V_O = (4/3)\pi r_O^3$, where r_O is the oxygen radius, and c_O is the atomic fraction of oxygen] (i.e., assuming oxygen atoms are perfect spheres). r_O of GeO_2 glass at 22.6 and 37.9 GPa and crystalline GeO_2 with CN = 6 are calculated by assuming an octahedral geometry ($r_O = r_{\text{GeO}}/\sqrt{2}$) in the same way as ref. 7. For GeO_2 glass with CN > 6, similar to the linear dependence of r_O between four- and sixfold-coordinated structures assumed in ref. 7, OPF is calculated by assuming a linear change of r_O between sixfold-coordinated structure and the ninefold-coordinated cotunnite-type structure, which is the next higher-pressure form in crystalline GeO_2 having a uniform CN > 6, predicted by first-principles calculations (28). To estimate the oxygen radius for the possible CN = 9 structure motif of GeO_2 glass, we first calculate the average O-O distance ($\langle \text{O-O} \rangle$) of the crystalline cotunnite-type GeO_2 , which consists of 21 O-O bonds at 1.929 (×2), 1.940 (×4), 1.948 (×4), 1.991 (×2), 2.055 (×2), 2.161 (×2), 2.243 (×2), and 2.540 (×3) Å, respectively (28). The oxygen radius r_O for cotunnite-type GeO_2 is then simply $r_O = \langle \text{O-O} \rangle/2$. Uncertainty of this r_O is assumed to be 1σ SD of $\langle \text{O-O} \rangle/2$. Uncertainty in OPF is estimated based on error propagation. For estimating r_O of GeO_2 glass with CN > 6, we calculated the r_O/r_{GeO} ratio (0.603) of crystalline cotunnite-type structure using its average Ge-O distance (1.735 Å) (28), and then, r_O of GeO_2 glass with CN = 9 structure motif is estimated from the measured Ge-O distances of GeO_2 glass at 49.4–91.7 GPa (Fig. 4A) by using $r_O/r_{\text{GeO}} = 0.603$. Uncertainties in the r_O/r_{GeO} ratio due to 1σ SD of the cotunnite-type GeO_2 are adopted as errors in r_O and the resultant OPF of GeO_2 glass with CN > 6.

Fig. 6 shows CN as a function of OPF extended to CN > 6, combined with the data of GeO_2 glass for $4 < \text{CN} < 6$ (7). Ref. 7 showed that fourfold-coordinated GeO_2 glass is stable in a wide range of OPF up to ~ 0.60 (Fig. 6). CN begins increasing when OPF approaches ~ 0.60 . The positive correlation between CN and OPF continues until OPF reaches ~ 0.64 . Then CN sharply increases to 6, with OPF remaining essentially constant at ~ 0.64 . At pressures of 22.6 and 37.9 GPa, OPF of GeO_2 glass increases, with CN remaining almost constant at 6 (Fig. 6). This is a stable plateau for sixfold-coordinated structure, similar to the plateau for CN = 4. The sixfold-coordinated GeO_2 glass is structurally stable in a wide range of OPF between ~ 0.64 and ~ 0.71 , similar to the behavior of sixfold-coordinated SiO_2 glass (7). Then, the CN of GeO_2 glass increases to 6.4 at OPF = 0.72. At higher pressures, CN increases further to CN = 7.4 at 94 GPa, while OPF remains essentially constant at ~ 0.72 (Fig. 6).

Table 1. Experimental pressure conditions and the results of the first (r1) and second peak positions of $G(r)$, and CN

Pressure (GPa)	r1, Å	r2s, Å	r2, Å	CN
0.0	1.738 ± 0.001		3.177 ± 0.01	4.0 ± 0.2
22.6 ± 0.4	1.846 ± 0.003	2.823 ± 0.02	3.209 ± 0.01	5.8 ± 0.2
37.9 ± 1.0	1.820 ± 0.001	2.795 ± 0.02	3.245 ± 0.02	5.9 ± 0.2
49.4 ± 1.3	1.824 ± 0.001	2.728 ± 0.02	3.149 ± 0.03	6.4 ± 0.2
61.4 ± 0.9	1.833 ± 0.001	2.734 ± 0.03	3.130 ± 0.03	7.0 ± 0.2
72.5 ± 1.1	1.814 ± 0.001		3.033 ± 0.12	7.0 ± 0.2
80.4 ± 0.3	1.813 ± 0.001		3.011 ± 0.01	7.1 ± 0.2
91.7 ± 0.6	1.810 ± 0.001		2.937 ± 0.06	7.4 ± 0.2

The second peaks at 22.6–61.4 GPa are fitted by two peaks for shoulder peak at ~ 2.7 – 2.8 Å (r2s) and main peak (r2).

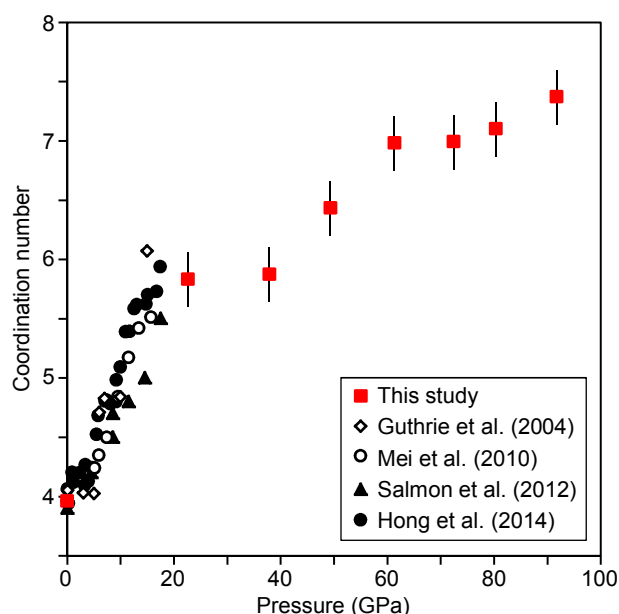


Fig. 5. Coordination number of Ge in GeO_2 glass as a function of pressure. Red squares represent results of this study, and black symbols are results of previous studies [open diamonds (1), open circles (3), solid triangles (27), solid circles (5)]. Coordination number at 22.6 and 37.9 GPa is almost constant at around 6, whereas it increases markedly to 6.4 at 49.4 GPa. Coordination number continues to increase with pressure, and the highest coordination number of 7.4 is observed at 91.7 GPa. Vertical bars represent errors of coordination number. Error of pressure is represented by the size of the symbols.

Ref. 7 has shown that the CN and OPF relationship is “universal” in many network-forming glasses and liquids such as SiO_2 , GeO_2 , silicate, aluminate, and germanate systems at high pressures in the CN range between 3 and 6. For example, for SiO_2 glass, the fourfold-coordinated structure is stable at OPF up to ~ 0.55 – 0.60 (7). At $\text{OPF} > \sim 0.60$, CN of SiO_2 glass increases to 6, and the sixfold-coordinated structure is stable at the OPF up to ~ 0.72 (7). The CN–OPF relation for SiO_2 coincides with that of GeO_2 as illustrated by the gray band in Fig. 6. Although experimentally determined highest CN in SiO_2 glass is 6 at 101.5 GPa (4), similar to crystalline GeO_2 , a cotunnite-type structure is also known as the next high-pressure form for crystalline SiO_2 having a uniform CN of 9 (29, 30). OPF of the cotunnite-type SiO_2 (30), calculated using the same method as GeO_2 , is 0.71, similar to that of cotunnite-type GeO_2 . Thus, similar to GeO_2 glass, SiO_2 glass may also change structure toward CN = 9 at higher pressure conditions. In addition to SiO_2 glass, the relation between CN and OPF in basalt melt also follows the same curve with CN between 4 and 6 (7), further confirming that the CN–OPF relationship describes a master curve for predicting structural changes in glasses and liquids at high pressure.

Our data on GeO_2 show that structural changes in glasses and liquids are closely associated with hard-sphere packing even to higher coordination numbers. Fourfold-coordinated structures are stable with OPF up to 0.55–0.60, corresponding to random loose packing of hard spheres (8, 9). Further compression increases CN sharply to 6 at OPF of ~ 0.64 , which is the most-disordered random closed-packing limit of hard spheres (8, 9), known as the maximally random jammed (MRJ) state (29) (Fig. 6). Our data show that there is a stable plateau for the sixfold-coordinated structure at OPF between ~ 0.64 and ~ 0.72 . OPF of ~ 0.72 is close to the maximal dense-packing state [the Kepler conjecture (KC) = ~ 0.74 , ref. 31], where a certain degree of local ordering, relative to the MRJ state, must be involved (32). Structure

evolves to higher CN with OPF remaining essentially constant just below the KC state packing limit (~ 0.74).

This extended CN versus OPF relation provides new insights into structural changes in other glasses and liquids under extreme compression. For SiO_2 glass, experimentally determined highest CN is 6 at 101.5 GPa (4). The number density ($0.154/\text{\AA}^{-3}$) and Si–O distance (1.67\AA) at 101.5 GPa (4) yield $\text{OPF} = 0.71$, which is still lower than that of the KC packing state. OPF of SiO_2 glass with CN = 6, calculated based on the results of ref. 4, increases from ~ 0.68 at 35.2 GPa to ~ 0.71 at 101.5 GPa (Fig. S1). If we linearly extrapolate the OPF–pressure trend, OPF of SiO_2 glass reaches 0.74 around 108 GPa, where structural change to CN higher than 6 is expected. We expect that further compression of silica (and likely other silicate glasses and liquids) should follow the dashed gray band shown in Fig. 6 toward higher CN.

Methods

The new double-stage large-volume cell is developed in a 200-ton PE press. We used a cup-shaped anvil with 12-mm cup diameter with 3-mm flat bottom as the first-stage anvil and (100)-oriented single-crystal diamond with 0.8-mm culet as the second-stage anvil. The gasket is composed of a cubic boron nitride + epoxy (10:1 in weight ratio) inner gasket with an aluminum alloy (7075) outer gasket. Initial sample size is 0.3 mm in diameter and 0.15 mm thick. A GeO_2 glass sample was packed in the gasket hole without pressure medium to avoid diffraction peaks from pressure medium in the X-ray diffraction measurement. A piece of Au (cut from 0.05-mm-diameter wire) is placed as a pressure standard at the sample edge to avoid contamination of X-ray diffraction signal from Au into X-ray spectrum of GeO_2 glass sample. Pressures were determined by the equation of state of Au (33). We measured pressure before and after structure measurement of glass because of long measurement time (~ 4.5 h). Pressure difference before and after structure measurement was up to 2.5 GPa. Table 1 shows the average pressures obtained before and after measurements with the pressure differences as errors. Pressure gradient between the center and edge of sample was up to 4 GPa at pressures up to 93.4 GPa obtained in a separate experiment using MgO and Au as samples.

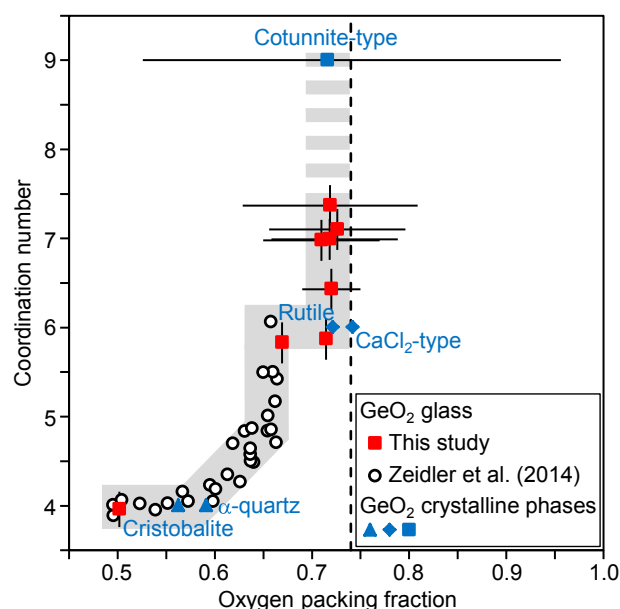


Fig. 6. Relationship between the OPF and the coordination number of Ge. GeO_2 glass data from this study (solid red squares) and from ref. 7 (open black circles) are compared with those of GeO_2 crystals with coordination numbers of 4 (cristobalite and alpha quartz, ref. 7) (blue triangles), 6 (rutile and CaCl_2 structures, ref. 21) (blue diamonds), and 9 (cotunnite-type structure, ref. 28) (blue square). OPF was calculated by using the method of ref. 7. Vertical broken line represents the maximal dense-packing state (KC = ~ 0.74 , ref. 31).

Structure of GeO_2 glass was investigated using the multiangle energy-dispersive X-ray diffraction technique (18). We used unfocused white X-rays. The size of incident white X-ray is 0.1 mm in both horizontal and vertical directions. We collected series of energy-dispersive X-ray diffraction patterns at 2θ angles of 4° , 5° , 7° , 9° , 12° , 16° , 21° , and 29° using a Ge solid-state detector (Canberra). Total exposure time to obtain the X-ray diffraction patterns for eight 2θ angles was about 4.5 h. Structure factor was derived from the observed energy-dispersive X-ray diffraction patterns using the program aEDXD (18). The Kaplow-type correction using an optimization procedure (34) was applied in determining final structure factor and pair distribution function. The iteration in the optimization process is typically 3.

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