Pressure-induced amorphization and existence of molecular and polymeric amorphous forms in dense SO$_2$

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We report here the pressure-induced amorphization and reversible structural transformation between two amorphous forms of SO$_2$: molecular-amorphous and polymeric-amorphous, with the transition found at 26 GPa over a broad temperature regime, 77 - 300 K. The transformation was observed by both Raman spectroscopy and x-ray diffraction in a diamond anvil cell. The results were corroborated by ab initio molecular dynamics simulations, where both forward and reverse transitions were detected, opening a window to detailed analysis of the respective local structures. The high-pressure polymeric-amorphous form was found to consist mainly of disordered polymeric chains made of 3-coordinated sulfur atoms connected via oxygen atoms, with few residual intact molecules. This study provides an example of polyamorphism in a system consisting of simple molecules with multiple bonds.

Polyamorphism is the counterpart of polymorphism observed in crystalline solids. It is characterized by the existence of two or more disordered forms, either amorphous or liquid, differing in local structural order whilst preserving the stoichiometry. This phenomenon is often accompanied by changes in coordination and density (1, 2) (for recent reviews see Refs. (3, 4)). Transformations between these different amorphous forms can be driven by pressure and temperature. While in the case of crystalline polymorphs the structural transitions are often (at least in principle) of first order and sharp, transitions in amorphous systems occur among isotropic forms and are more likely to be continuous. This is related to the absence of constraints prescribed by lattice periodicity, allowing for a gradual transformation between very different forms.

The first and perhaps most celebrated example of polyamorphic behavior was discovered in water ice by Mishima in 1984 (5, 6), observing that compression of ice I$_h$ at 77 K induced a transformation to an amorphous state. Through specific compression/decompression/heating protocols, at least two different forms of amorphous water ice were identified, known as low-density amorphous (LDA) and high-density amorphous (HDA) ices. The local structural order in the HDA and LDA ices differs by the presence of nonbonded water molecules in the first coordination shell of the former, resulting in the two forms having substantially different density. The existence of two amorphous forms of water ice was suggested to be related to existence of liquid-liquid transition and a second critical point of water (7).

Since, similar phenomena have also been observed in other systems such as Si (8), SiO$_2$ (9–11), GeO$_2$ (12), where the polyamorphism is related to a change from tetrahedral to octahedral coordination at high pressure. Other examples where pressure induces changes in simple molecular systems include amorphous S(13) and liquid Si(14) (for more examples and review see Refs. (3, 4)). Dramatic structural changes leading to amorphization have been observed upon compression of molecular crystals where multiple bonds are present. It is well known that pressure can destabilize multiple bonds in molecules in favour of extended polymeric networks with a higher coordination. Due to the associated strong kinetic effects, creation of amorphous phases are often observed, especially when compression is performed at low temperatures. Amorphization of molecular crystals at high pressure have been observed in the famous examples of nitrogen (15, 16),

Polyamorphism | sulfur dioxide | high pressure | polymeric form

Significance Statement

Some substances are known to exist in several different structurally disordered solid states and transform between them similarly to structural phase transitions between crystalline polymorphs. This interesting and yet not fully understood phenomenon is called polyamorphism and notable examples include water, SiO$_2$, Si, etc. Here we present a new example of such behavior in a simple molecular substance, SO$_2$. By using experimental high-pressure techniques we observe a reversible transition between different amorphous states around 26 GPa. Experimental results are well supported by ab initio simulations, which identify the high-pressure amorphous form as non-molecular, polymeric one consisting of intertwined chains. Our findings contribute to the fundamental understanding of structure of disordered matter as well as high-pressure behaviour of simple archetypal molecules.

H.C.Z. performed the experiments and contributed to the data analysis. O.T. performed simulations, analyzed the data and wrote the paper. X.D.L., R.B. and E.G. contributed to the discussion of results and wrote the paper. X.D.L., M.S. and F.A.G. conducted the experiments and wrote the paper. P.D.S. and E.G. contributed to the discussion of results and wrote the paper. M.S. and F.A.G. designed the research, contributed to the plan the experiments, to the data analysis and wrote the paper. R.M. designed the research, contributed to the analysis and discussion of results and wrote the paper.

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carbon dioxide (17, 18) and benzene (19–21). In nitrogen, the strong triple bond of the molecule breaks under high pressures giving rise to a single-bonded network. Whilst in carbon dioxide, the double bond becomes unstable and carbon coordination increases to 3 and 4. In the case of benzene the aromatic ring opens and a network of hydrogenated carbons with single bonds is formed. The parent crystalline states of these amorphous materials have been discovered both in nitrogen (22, 23) and in carbon dioxide (24, 25) after high temperature annealing obtained by means of laser heating.

Sulphur dioxide is an important molecule in chemistry, serves a significant role in industrial applications, and has been attributed to atmospheric and geological processes. Unlike CO2, the SO2 molecule is bent, described by two resonant structures with one single and one double bond (26, 27). The crystalline forms of SO2 have been previously experimentally studied at pressures up to 32 GPa by Song et al. (28). Here, we present a combined experimental and computational study of SO2 up to pressures of 60 GPa and over a broad temperature regime. Observations via Raman spectroscopy and X-ray diffraction, well supported by ab initio simulations, provide a detailed description on the atomistic level of the transformations under compression/decompression cycles. These findings give evidence of a hitherto unobserved example of polyamorphism related to a reversible transformation between molecular and polymeric amorphous forms of SO2.

In Fig. 1, we present a selection of Raman spectra measured upon increasing pressure up to 60 GPa, at 77 K (panel A), as well as subsequent decompression back to ambient pressure at room temperature (panel B). Similar experiments for compressions at 210 K were also conducted and reported in Fig. S1 Supp. Mat. We find in general, an agreement between our Raman measurements of solid molecular SO2 under pressure and those reported previously (28). However, the dramatic spectral changes at low temperatures were not observed previously as pressures were limited to only 22 GPa (28). From these datasets, what is readily evident is the progression from sharp molecular peaks of SO2, i.e., $\nu_1$ 1050-1220 cm$^{-1}$, $\nu_3$ 1240-1320 cm$^{-1}$, and $\nu_2$ 520-600 cm$^{-1}$, to much broader, weaker peaks upon compression. In addition, new broad and weak bands appear at different frequencies, indicating pressure-induced amorphization together with changes in the local structure. Upon decompression, the sharp molecular peaks of SO2, $\nu_3$, $\nu_2$, and $\nu_1$, are recovered while the new broad bands disappear at the same time, showing that amorphization and overall changes in the local structure are reversible. Panel C: evolution of vibrational density of states (VDOS) from ab initio MD simulations along compression (solid lines) and decompression (dashed lines). Color represents structural state of the system: black - molecular crystal, red - molecular amorphous and blue - polymeric amorphous. Star in panel B marks ruby peaks.

SO2 undergoes a structural transformation into an amorphous form whilst preserving its molecularity, i.e., consisting of SO2 molecular units. Amorphization could have been enhanced by the shear stress, which is in turn related to the deformation of the gasket hole. This is supported also by our DFT calculations (see section on simulation results). Upon further compression, above 22-25 GPa, additional and significant modifications to the Raman signature are observed, the $\nu_2$ and $\nu_3$ peaks progressively diminish and are barely observable above 30-34 GPa. Conversely, the strongest Raman excitation of SO2, $\nu_1$ (1170-1200 cm$^{-1}$ (Fig. 1 A)), is visible to the maximum pressures of 50-60 GPa (see also Fig. 1 B), albeit comparatively weaker. The $\nu_3$ band peak is found to merge with an altogether new weak and broad peak centred at 1220-1230 cm$^{-1}$, denoted as $\alpha$ in Fig. 1 A. In addition to emergence of the $\alpha$-peak, another new and broad band appears, at 600-1000 cm$^{-1}$ with
a high frequency edge at around 900 cm\(^{-1}\), which we call \(\beta\). Both bands \(\alpha\) and \(\beta\) appear to be of non-molecular origin, which suggests that the emergence of these excitations signals that amorphous-molecular \(\text{SO}_2\) undergoes a transformation into a non-molecular/extended amorphous form.

Comparison with the pressure-induced molecular to amorphous-non-molecular transformation in \(\text{CO}_2\) (17, 18, 25) can help to interpret the transformation observed here in \(\text{SO}_2\), as it bears similarities. Carbonia, the non-molecular amorphous \(\text{CO}_2\), has been shown to be made of a mixture of \(C\) in a 3-fold and 4-fold oxygen coordination in similar proportions. The 3-fold coordinated \(C\) sites are uniquely identified by \(C=O\) stretching peaks in the Raman and the IR spectrum, 1900-2000 cm\(^{-1}\) at 50-60 GPa, which roughly corresponds to the average value, \(\nu\) (stretch \(\text{CO}_2\)), between the symmetric and the antisymmetric stretching modes of molecular \(\text{CO}_2\). However, when considering the full Raman and IR spectra, the single \(C=O\) bond stretching and deformation modes ascribed to both 3-fold and 4-fold coordinated \(C\) sites, form a broad spectral distribution extending over 500-1500 cm\(^{-1}\), approximately 0.26-0.77 of the average \(\nu\) (stretch \(\text{CO}_2\)). Consequently, in the case of \(\text{SO}_2\), the aforementioned peak, \(\alpha\), is roughly the average frequency of the two molecular stretching modes, and can therefore be attributed to the \(S=O\) stretching modes for non-molecular/extended \(\text{SO}_2\) with \(S\) in 3-fold coordination by \(O\). In addition, in the frequency range given by 0.26-0.77 of \(\alpha\), 320-940 cm\(^{-1}\), we observe the previously described band \(\beta\). Therefore, again in accordance with the \(\text{CO}_2\) analogue, the \(\beta\) band can be attributed to single \(S=O\) bond stretching and deformation modes in the non-molecular/extended \(\text{SO}_2\). A discrepancy of using carbonia as an analogue however is that the 4-fold oxygen coordination is absent in \(\text{SO}_2\), in accordance with our DFT simulations (discussed later). We do however also have a two component system, considering that the \(\nu_1\) peak for molecular \(\text{SO}_2\) is still present at the highest pressures, although weak. Therefore, inferring that the overall non-molecular/extended \(\text{SO}_2\) consists of a mixture of trace 2-fold \(S\) sites, still molecular in nature, in a bulk of non-molecular 3-fold coordinated \(S\) sites. An alternative possibility compatible with experimental data would be that molecular parts of the sample with 2-fold coordinated \(S\) and non-molecular parts with \(S\) in higher coordination are phase separated on a macroscopic/mesoscopic scale.

On decompression, the spectral changes and transformations identified previously on compression are reverted, demonstrated in Fig. 1 B. However, a minor hysteresis is observed, attributed to the kinetically limited structural changes. We observe that when the sample is decompressed below 30-25 GPa, both the \(\alpha\) peak and \(\beta\) band disappear, whilst the well-defined molecular peaks \(\nu_1, \nu_2,\) and \(\nu_3\) emerge suddenly. Further, in the low frequency/lattice region below 350 cm\(^{-1}\) a diffuse, liquid-like band clearly develops, with no additional substantial changes down to about 5 GPa. These changes demonstrate again a change in the amorphous structure, recovering molecular \(\text{SO}_2\) units at 25-5 GPa on decompression. This molecular-amorphous phase is then found to further transform into a crystalline molecular \(\text{SO}_2\) structure below 5 GPa, indicated by the sharp lattice mode peaks observed below this pressure and crystalline x-ray diffraction (discussed later). Interestingly, an additional peak is found below 20 GPa at around 600 cm\(^{-1}\) which was also observed in a previous study, where it was attributed to the formation of molecular clusters (28). The reversible transformations of \(\text{SO}_2\) to molecular and non-molecular amorphous forms again parallel the similar case of \(\text{CO}_2\) (17, 18, 25), and at odds with the cases of aromatic molecules, where instead amorphous non-molecular forms obtained at high pressures are recoverable to ambient conditions (19–21, 29, 30).

The non-crystalline nature of \(\text{SO}_2\) at high pressure has also been assessed by x-ray diffraction, and the evolution of the static structure factor upon room temperature decompression of sample compressed at low temperature is shown in Fig. 2 A. The static structure factor has been obtained by the empty cell subtraction, taking into account the form factors of oxygen and sulfur as well as the Compton contribution from the sample, following a procedure described elsewhere (31). In general, it is found that at the highest pressures the two peaks at about 2.2 and 3 Å\(^{-1}\) are of similar intensity, while on decompression the first peak significantly increases with respect to the second one which at the same time moves to lower \(Q\).
comparison between patterns measured at 35.5 and 23.5 GPa, there is a clear change of the static structure factor. The first contribution, centred at 2.2 Å⁻¹, becomes more prominent and sharpens markedly, whilst the second has a more subtle loss of intensity. These changes, corresponding to the transformation from an extended amorphous to a molecular amorphous form, are in strong agreement with the Raman results outlined previously.

In summary, we have observed two distinct structural transformations. At lower pressures, the transformation from a molecular crystal to a molecular, van der Waals type, amorphous solid, with no changes in the molecular unit and to a non-molecular/extended state on further compression. Interestingly, a substantial effect of temperature is observed for the former transformation, despite the expected energy barrier to be comparatively small. For example, instead of an expected isobaric phase line, the transformation occurs at 5 GPa lower pressures at 210 K compared with 77 K isotherm. Conversely, the second transformation, to a non-molecular/extended amorphous state, is a chemically reconstructive one, and therefore would logically have a higher associated energy barrier and consequently a stronger temperature dependence. Instead, this transformation is observed at roughly the same pressures for both isotherms studied, 77 K and 210 K, at around 25-30 GPa, counterintuitively suggesting that the barrier for polymerization is small in relation to the initial molecular amorphisation. We note that this is quite different from the case of CO₂ and N₂, where a much stronger hysteresis is reported (23–25).

In order to obtain a better understanding of the processes on the atomistic level, we performed ab initio MD simulations following a pressure path akin to the experiment. We first performed a test in order to check whether applying shear stress to a perfect Aea2 molecular crystal at low pressure might result in amorphous molecular structure as observed in experiments. We gradually induced shear strain by deforming the γ angle of the supercell by up to 30° and observed transformation into a disordered molecular form, confirming the experimentally observed amorphization.

The full simulation protocol is shown in Fig. S3 in Supp. Mat. In order to start the compression from a well-defined amorphous molecular structure, we melted a perfect Aea2 molecular crystal (32) in a 3 × 3 × 3 supercell (108 SO₂ molecules, equivalent to a 324 atom unit cell) by heating at P = 0 GPa to 600 K. Through subsequent cooling to 0 K, we prepared the amorphous structure, which served as an initial configuration for further simulations. Following the experimental pathway, we performed a gradual compression to 60 GPa and subsequent decompression to 10 GPa (in 10 GPa steps) at 300 K in order to accelerate the structural transformations (both on compression and decompression). Interestingly, at this temperature we observed some diffusion of molecules in the molecular phase, suggesting that the sample might possibly be in metastable liquid regime.

Analysis of the partial radial distribution functions (RDFs) (Fig. 3), obtained from simulations, can provide a more detailed description of the observed transformations on an atomistic level. The RDFs clearly indicate the reversible amorphous to amorphous transformation, corresponding to the S in 2- to 3-fold O coordination change, providing further evidence for the experimentally observed phases and their associated reversibility. Upon compression of the initial molecular amorphous sample to 10 GPa at 300 K, SO₂ retains its molecular units. Evidenced in Fig. 3 A, the peak at 1.44 Å, corresponding to the double bond, is sharp and well separated from the next neighbor at 2.5-3.0 Å and the coordination number of S atoms with respect to O atoms is 2. Upon compression to 20 GPa modifications in the RDF are observed, and at 30 GPa there is a substantial change, the first peak becomes weaker whilst a new peak at slightly longer distance, 1.6 Å, appears. These changes indicate that some of S=O double bonds are
broken and replaced by single ones. In the same pressure regime, 3-coordinated S-atoms appear with 2 single S-O bonds and one S=O double bond, forming polymeric chains (see snapshots shown in Fig. 4). On further compression, the first peak is found to progressively diminish in intensity whilst the second is enhanced, resulting in 82% of S atoms to be in a 3-fold oxygen coordination at 60 GPa. On decompression (see Fig. 3 B), akin to experimental observations, we observed the reverse evolution, further demonstrated by the pressure dependence of the number of 2- and 3-coordinated S atoms in Fig. 3 C. Additionally, at 10 GPa the previously identified polymeric chains observed on compression disappear entirely (see Fig. 4), and the system reverts back to its initial molecular amorphous state. The strong agreement between experiment and simulations, means that insights from the MD-calculations can identify the experimental observations as forward and backward transitions between molecular and polymeric amorphous forms of SO$_2$. The dependence of the coordination number of S atoms on pressure (Fig. 3 C) exhibits some hysteresis, which suggests that the transition might have a weakly first-order character. The pressure dependence of density upon compression and decompression, shown in Fig. S4 (Supp. Mat.) shows a very small hysteresis and does not exhibit any particular features across the structural transformation. We performed a similar compression simulation also at 500 K and found the polymerization to start at 30 GPa, suggesting that the transition is shifted to higher pressure upon increasing temperature.

Above 20 GPa, the calculated static structure factor, $S(Q)$, shows important changes upon compression (Fig. S2 Supp. Mat.). The intensity of the first diffraction peak at about 2.2 Å$^{-1}$ drops, while a new peak appears, around 3 Å$^{-1}$, which grows with increasing pressure. At 60 GPa, the height of both peaks becomes similar, reflecting XRD patterns above 35.5 GPa in Fig. 2 A. All changes are found to be reversible upon decompression, albeit with a small hysteresis, and the calculated structure factor agrees very well with the experimental one (Fig. 2). Simulations allow us to decompose the total $S(Q)$ into contributions from atomic pairs (Figs. S5 Supp. Mat.). The first peak, around 2.2 Å$^{-1}$, originates mainly from non-bonded S···S pairs, while the weaker second peak, at 3 Å$^{-1}$, comes from O···O and finally the broad peak, around 5 Å$^{-1}$, is mainly due to the S-O pairs. It is clear that the loss of the first peak of total $S(Q)$ is a consequence of changes in the S-O contribution, which exhibits a pronounced drop above 20 GPa (Fig. S5 Supp. Mat.). Therefore, the observed evolution of $S(Q)$ can directly reflect changes of the distance of S-O neighbors upon polymerization.

On compression, comparing the vibrational spectra from simulations (Fig. 1 C) with experimental Raman spectra (Fig. 1 A), we observe qualitatively similar evolution. Above 20 GPa the distinct $\nu_2$ peak at 550 cm$^{-1}$ progressively disappears whilst the 400 - 500 and 600 - 900 cm$^{-1}$ regions become enhanced. Meanwhile, the two molecular peaks, $\nu_1$ around 1100 cm$^{-1}$ and $\nu_2$ above 1200 cm$^{-1}$ gradually merge into a single broad peak around 1200 cm$^{-1}$, again in agreement with experiments (Fig. 1 A). The evolution of the peaks can be understood from the projected VDOS, allowing decomposition of the total VDOS into contributions from structurally distinct S and O atoms (Fig. S6 Supp. Mat.). Both S and O atoms can be either in molecules or in polymeric chains. Moreover, O atoms in polymeric chains are either at a bridging position between two S atoms (S-O-S) or at terminal position, doubly bonded to S atoms (S=O). The evolution of the total VDOS clearly reflects the gradual conversion of molecules into polymeric chains. Again, upon decompression the reverse evolution is observed, in agreement with experiment (Fig. 1 B).

In order to understand the origin of the two amorphous phases it is useful to discuss their underlying crystalline counterparts. SO$_2$ forms at ambient pressure and low temperature below 201 K, an $\text{Aea}_2$ molecular crystal(32). Instead, its analogue SeO$_2$ forms at ambient pressure a $\text{P4}_2/mbc$ polymeric crystal consisting of chains (mineral Downeyite (33)). The possible existence of polymeric SO$_2$ (polysulphite) was studied in Refs.(34, 35) where energies of various oligomers were calculated. Additionally, in Ref.(35) a crystal structure with infinite polymeric chains was studied obtained by substituting Se atoms in Downeyite by S atoms. It was concluded that polymeric SO$_2$ is energetically higher than its molecular form and it was proposed that the polymer could be stabilized at
high pressure. This can be understood by applying the well-known pressure-homology rule (36), (37), (38) stating that light elements behave at high pressure like more heavy elements from the same group at lower pressures.

The structure of polymeric phases of SO₂ has not been, to our knowledge, determined experimentally. In order to check its possible existence at high pressure we performed a structural search of crystalline phases of SO₂ employing evolutionary approach (details are described in Methods). We show here the main results of our search. The enthalpy vs. pressure graph for low-enthalpy phases is shown in Supp. Mat. Fig. S7. At zero pressure we found the molecular crystal structure with space group $Aea_2$, in agreement with experiment (32). Upon increasing pressure we found another molecular crystal structure $P$$\overline{2}$$\overline{1}$/c which becomes stable above 1 GPa. Besides this structure there are two low-lying metastable molecular structures $Pc$, and $Cc$, which around 11 GPa transform into different molecular forms with space groups $Pmc_2_1$ and $Ama_2$, respectively and become stable (both have very similar enthalpies) with respect to the $P$$\overline{2}$$\overline{1}$/c one. Upon further compression they transform into polymeric structures (all structures are shown in Supp. Mat. Fig. S8 and cif files of the $Pmc_2_1$ and $Ama_2$ structures are included in Supp. Mat.). Both polymeric structures still have very similar enthalpy and conformation of chains and differ only in chain stacking. The mechanism of polymerization is illustrated in Fig. S9 Supp. Mat. showing the pressure evolution of bond-lengths upon compression. A similar polymerization mechanism is likely to apply also in the amorphous phase when molecules of suitable orientation approach each other. Both polymeric structures have the same conformation of chains as SO₂ downeytne but the stacking of chains is different. The stable polymeric structure has at 20 GPa bond length of 1.46 Å between S atom and terminal oxygen (S=O) and 1.65 Å between S atom and bridging oxygen (S-O), similar to the peak positions of S-O RDF (Fig. 3). The sulphur atom in the chain is surrounded by three oxygen atoms in roughly trigonal pyramid coordination (at 20 GPa bond angle O-S-O is 90° and O=S=O is 100°) suggesting presence of sp² hybridization (see also Ref. (35) where it was suggested that the dimerization of SO₂ is related to sp² → sp³ rehybridization). In order to quantitatively assess the bond order of S-O bonds in molecular and polymeric SO₂ we employed the DDEC6 atomic population analysis method (39).

We found a bond order of 2.12 in molecule and 2.14 and 1.15 for S=O and S-O bonds in the polymeric phase. In Fig. S10 Supp. Mat. we show the e-DOS including projections on $s$ and $p$ orbitals for the crystalline molecular and polymeric phase (no $d$-orbital participation was found). We also note that the diffraction pattern of the recrystallized phase upon decompression to 6.5 GPa (Fig.2 A) shows a strong similarity with the diffraction pattern of the molecular phase $Aea_2$ at the same pressure (for comparison see Fig. S11 Supp. Mat.). This provides an additional evidence that the system after the compression/decompression cycle returns to its parent state.

To assess quantitatively the effects of structural disorder, we calculated the enthalpy of the amorphous phase (relaxed to $T = 0$) and found it at 10 GPa to be about 0.26 eV/formula unit above the crystalline phases (it is shown in Fig. S7 in Supp. Mat.). We note that because of the short time scale available in the ab initio MD simulations (of the order of 10 ps) the amorphous structure might not be fully relaxed structurally. We also determined with our DFT simulations the electronic properties of the a-SO₂ phase. The system does not metalize up to 60 GPa, with band gap of at least 0.6 eV in PBE approximation.

**Conclusions and outlook.** We observed a pressure-induced amorphization and a reversible structural transition between molecular and polymeric amorphous forms of SO₂ at pressures around 26 GPa. The transition has small hysteresis pointing to the fact that the associated kinetic barriers are low. The lower pressure of the transition between molecular and polymeric amorphous forms, as well as the back transformation, is qualitatively facilitated by the molecular polarity. This, supported by the high density attainable under pressure, drives the intermolecular interaction and lowers the activation energy of the transformation. To our knowledge this kind of transition was not yet observed and provides a new example of structural transition between disordered non-equilibrium states of solid matter. Unlike in a-CO₂, where polymeric a-carbonia contains 3- as well as 4-coordinated C atoms, here the molecular form converts into polymeric form with only 3-coordinated S atoms. It will be of interest to study whether the two amorphous states continue to exist also in liquid state, either in stable or metastable (undercooled) region. Further experimental and theoretical work is necessary to accurately map the solid and liquid regions and uncover further details of the phase diagram of SO₂.

**Materials and Methods**

**Experimental methodology.** The SO₂ gas was loaded into the DAC by means of cryogenic loading; the gas was condensed between one diamond anvil and the gasket placed on the other diamond of a DAC which was opened by few mm and cooled to liquid nitrogen temperature inside a sealed glove box purged with nitrogen to avoid moisture condensation. We performed Raman spectroscopy using a state of the art confocal Raman microscope with 15 and 2 micron of axial and transverse resolution respectively. The spectrometer consisted of a Spectra Pro 750 mm monochromator, equipped with Pixis Princeton Instrument CCD detector. Bragg grate filters were used to attenuate the laser light and spatial filtering of the collected light to obtain high quality spectra down to 7 cm⁻¹ with minimal background from the diamond anvils and strong signal from the sample. The laser beam was expanded and cleaned by a Band Pass Filter. We used a Laser Taurus at 660 nm with 10 mW of power and Laser Ventus at 532 nm with 0.5 mW of power to check for the presence of eventual fluorescence bands in the spectrum. We generally used a 300 gr/mm grating as the spectral features were getting very broad and weak with pressure. The pressure was determined by the fluorescence of a small ruby placed in the sample or from the diamond stressed edge which we detect with high accuracy thanks to the excellent spatial resolution of the setup. The XRD measurements were made at Petra (proposal ID: I-20181128) using monochromatic X-ray beam with 42.7 keV energy ($\lambda = 0.2922$ Å) and the scattered X-rays were detected by a Perkin Elmer XRD1621 (2048x2048 pixels, 200 x 200μm) detector. The diffraction patterns have been measured only along decompression of an amorphous sample obtained from a compression at low temperature while monitoring the changes with Raman spectroscopy. The excellent transverse spatial resolution allows to obtain clean diffraction patterns of the sample without the presence of spurious diffraction lines from the metallic gasket. The empty cell subtraction, which is of fundamental importance to obtain reliable measurements of the diffuse scattering from an amorphous or liquid sample in the DAC, has been in this case easily obtained by measuring the empty cell at the end of the decompression run when the SO₂ has completely back transformed to the gas state and escaped from the sample chamber.
Simulations methodology. We performed a structural search for crystalline phases of SO₂ employing the USPEX (40) package at pressures 10, 20 and 50 GPa with 4 formula units (12 atoms) within unit cell. Ab initio simulations were performed by density functional theory (DFT) as implemented in VASP 5.3 and 5.4 codes (41–43), employing projector-augmented-wave pseudopotentials (with 6 valence electrons for both S and O atoms) and Perdew-Burke-Ernzerhof (PBE) (44) parametrization of the GGA exchange-correlation functional. In evolutionary search, structural relaxations and enthalpy calculations we employed the harder Sρ and Oρ pseudopotentials with cutoff of 910 eV while in MD calculations we used the regular ones S and O with cutoff of 520 eV. Compression, decomposition and heating were performed by 6 ps variable cell NpT simulations with Langeniv thermostat and Τ-point Brillouin zone sampling. We used 10.0 and 2.0 ps⁻¹ friction coefficients for atomic and lattice degrees of freedom respectively, and 10000 mₐ/s as barest fictitious mass. Data for velocity autocorrelation function were generated by equilibrating sample for 6 ps in NpT and then running 20 ps NVE simulation. Total and projected vibrational density of states (VDOS) were computed in standard way as Fourier transform of mass-weighted velocity autocorrelation function from MD trajectories at pressures from 10 to 60 GPa. Static structure factors S(Q) were calculated by performing Fourier transform of the RDFS from MD trajectories at several pressures along compression and decompression runs.

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