Structure and SHG of the high pressure phase IV of HgBr$_2$

Marc Hostettler$^a$,*, Dieter Schwarzenbach$^b$, Jan Helbing$^c$, Vladimir Dmitriev$^d$, Hans-Peter Weber$^{b,d}$

$^a$Laboratorium für chemische und mineralogische Kristallographie, University of Bern, CH-3012 Bern, Switzerland
$^b$Laboratoire de cristallographie, EPFL/SB/IPMC/LCR, Swiss Federal Institute of Technology Lausanne, CH-1015 Lausanne, Switzerland
$^c$Institut de Physique de la Matière Condensée, University of Lausanne, BSP Dorigny, CH-1015 Lausanne, Switzerland
$^d$Swiss Norwegian Beam Lines, ESRF, BP 220, F-38043 Grenoble Cedex, France

Received 2 October 2003; accepted 5 November 2003 by P. Wachter

Abstract

Angle dispersive X-ray diffraction experiments on mercuric bromide (HgBr$_2$) under high pressure up to 11.0 GPa were carried out at room temperature using synchrotron radiation. In addition to the already known four different phases of HgBr$_2$ in the pressure–temperature range of $p < 4.5$ GPa, $90 < T < 600$ K our observations show the existence of a new phase (V) above 9.0 GPa and, together with published material, support the phase transition sequence: (I) orthorhombic–(II) orthorhombic–(III) monoclinic–(IV) trigonal–(V) trigonal/hexagonal. The structure of phase IV with space group symmetry $P3$ has been determined from powder diffraction data. The observation of second-harmonic-generation signals confirms the absence of an inversion center. The structure of phase IV is a commensurately modulated variant of the CdI$_2$ type layer structure, where part of the Hg atoms are displaced from the centers of the HgBr$_6$ octahedra by a much as 0.76 Å in the direction perpendicular to the layers.

$^*$ Corresponding author. Tel.: +41-21-631-4274; fax: +41-31-631-3996.
E-mail address: marc.hostettler@krist.unibe.ch (M. Hostettler).

PACS: 61.50.Ks; 81.30. – t; 42.70.Mp

Keywords: E. High pressure; E. Synchrotron radiation; C. Crystal structure and symmetry; D. Phase transitions

1. Introduction

The solid phases of the mercuric halides HgX$_2$ (X = I, Br, Cl) form two kinds of crystal structures: (i) closest sphere packings of X with Hg in tetrahedral interstices; (ii) linear or bent HgX$_2$ molecules resulting in distorted closest packings of X. HgI$_2$ is the most intensively investigated mercuric halide; its red modification is used as room-temperature detector material for X- and $\gamma$-rays [1–4]. Besides the stable red phase, HgI$_2$ also forms an orange and a yellow phase at ambient conditions, both of which are metastable [5]. The orange phase comprises three different closely related structures, which may occur together in a multiply twinned conglomerate [6]; the structure of the yellow phase [5] is related to the high-temperature yellow phase [7]. The structural motifs of these polymorphs are: corner-shared HgI$_4$ tetrahedra (structure kind i, red), corner-shared Hg$_6$I$_{10}$ supertetrahedra (i, orange), and HgI$_2$ molecules (ii, yellow).

At ambient conditions, phase I of HgBr$_2$ adopts the structure of the yellow metastable polymorph of HgI$_2$ [8,9]. It is orthorhombic, polar space group symmetry $Cmc_2_1$ ($D_2h$), and thus presents opto-electronic properties; it is also a candidate material for the detection of $\gamma$-rays [10]. At ambient pressure, phase I is stable from 90 K, the lowest temperature of observation, to the melting temperature of 509 K. The application of pressure up to 4.5 GPa induces several transitions as revealed by dilatometry (compression-curve method) [11] and Raman spectroscopy [12]. Three phases (II–IV) have been reported in this pressure range.
the structures of which have not yet been determined. In order to better understand the behavior of this material under pressure, we performed X-ray powder diffraction and second-harmonic-generation (SHG) experiments using diamond-anvil cells (DAC). In this communication, we present an overview of the diffraction results, the crystal structure of phase IV and its SHG signal, as well as the occurrence of an additional phase V.

2. Experimental

2.1. X-ray diffraction

In situ high-pressure angle-dispersive X-ray diffraction data were recorded at the Swiss–Norwegian Beam Lines at the European Synchrotron Radiation Facility (ESRF, Grenoble, France) by using monochromatic radiation with wavelength $\lambda = 0.7498(2)$ Å collimated to a diameter of 100 μm. Commercial HgBr$_2$ powder (Fluka, puriss. p.a. > 99%) composed of needle shaped crystallites was ground to an average particle size of 5–10 μm. The sample was introduced into the hole of a tungsten gasket in a DAC together with a ruby chip and silicone oil as pressure medium. The pressure was determined by the fluorescence of the ruby. Diffraction data were recorded with a MAR345 area detector at several pressures up to 11 GPa. The 2D images were corrected for geometrical distortions and transformed into a 1D powder diagram by integration with program Fit2D. Diffraction intensities were corrected for absorption of the DAC.

2.2. Second-harmonic generation

The first description of a SHG experiment using powders has been reported in Ref. [16]. We have performed a similar experiment using a femtosecond Ti:sapphire laser at 810 nm. Second-harmonic signals were measured in transmission geometry. Filters and a spectrometer were used to suppress the 810 nm light, and spectra in the 405 nm region were recorded with a CCD detector. Powdered sample contained in a capillary was first investigated at ambient conditions (phase I). The same experiment was repeated with a DAC, at a pressure of 6.2(1) GPa (phase IV) and signals were scaled to the respective sample volumes.

3. Results

3.1. Phase transition sequence

Up to the III–IV transition at 4 GPa, the powder rings show the sample to be strongly textured as evidenced by the raw 2D images of Fig. 2. The diagrams of phase II are very similar to those of phase I. Because of the texture, we could not identify any structural change between these two phases. The I–II transition was first detected by dilatometry. It is accompanied by a volume change similar to the one observed for the II–III and III–IV transitions [11]. However, its existence was not conclusively confirmed by Raman spectroscopy, the reported differences between I and II being minor [12]: for I, the observed peaks were at 16, 57 and 186 cm$^{-1}$, while for II the first peak was absent and the others had shifted by $+5$ and $-2$ cm$^{-1}$, respectively. Phase III displays a more complicated diffraction pattern, with additional powder lines, suggesting a lower, possibly monoclinic symmetry. However, also for this phase the texture did not permit us to propose a convincing indexation. In addition, the corresponding Raman spectrum shows inferior signal to noise compared to the spectra of phases I, II and IV [12]. These difficulties might be due to the formation of a strongly disordered state in the stability range of phase III of HgBr$_2$. At the III–IV transition the texture disappears. This
3.2. Second-harmonic generation

Both phases I and IV of HgBr₂ generate a clear second-harmonic signal. The observed integrated intensities are scaled with the respective volume of the sample to allow the comparison of the relative conversion efficiencies of the two phases, giving 250,000 and 230,000 photons s⁻¹ mm⁻³ for phase I and IV, respectively. These values are of the same order of magnitude as those measured for a powdered sample of the yellow high-temperature phase of HgI₂ (space group symmetry P2₁), and about three orders of magnitude larger than the value measured for the centrosymmetric red phase of HgI₂ at room temperature [7]. Red HgI₂ was used as a centrosymmetric reference material to estimate the magnitude of a possible surface effect for our experimental set-up. The structures of phase I of HgBr₂ and of the yellow high-temperature phase of HgI₂ are known to be non-centrosymmetric and the signal measured for phase IV is too large for a surface effect. Therefore, it is safe to conclude that the inversion is also absent in the structure of phase IV of HgBr₂.

3.3. Structure of phase IV

The X-ray diffraction intensities do not present any systematic absences. The structure was solved with direct methods [17]. Different hexagonal and trigonal space groups were tested and an initial model was obtained in the non-centrosymmetric space group P3, in agreement with the observed second-harmonic signal. This structure was refined with the Rietveld method using program GSAS [18], leading to the confidence factors: \( wR_p = 0.035 \), \( R_p = 0.025 \) and \( R(F^2) = 0.075 \) [19]. The experimental data, refined profile and their difference curve are shown in Fig. 3. The structural parameters are given in Table 1.

The structure of phase IV of HgBr₂ is a derivative of the structure of CdI₂. It is built from layers of edge-sharing octahedra that are stacked through Br·Br van der Waals contacts. In contrast to the CdI₂ type structure, some of the coordinated metal atoms are displaced along \( c \) from the centroids of the octahedra, leading to three distinct mercury sites and thus \( Z = 3 \) instead of \( Z = 1 \). Fig. 4 presents a view of the structure onto the plane of the layer \([0 0 1] \), and an edge view of the layer down the \([1 1 0] \) direction showing the alternating displacements along \( c \) of the three mercury atoms in neighboring octahedra along \([1 1 0] \). This structure may be described as a commensurate modulation of the CdI₂ structure. The Hg–Br bond lengths are 2.56(1), 2.60(1) Å for Hg₁, 2.60(1), 2.72(1) Å for Hg₂ and 2.63(2), 3.10(1) Å for Hg₃. Hg₁, Hg₂ and Hg₃ are displaced from the centers of the coordination octahedra by \( +0.02, -0.19 \) and \( +0.76 \) Å, respectively. Thus, Hg₃ adopts a trigonal coordination whereas Hg₁ and Hg₂ show an approximately octahedral coordination with six bonds of roughly the same length. The interlayer Br·Br contacts are 3.48(1), 3.64(2) and 3.78(2) Å. Surprisingly, the average isotropic displacement parameter of the Hg atoms is 1.7 times larger than the one of Br. Even if the atomic displacement parameters derived from powder data are often unrealistic, this observation of larger displacement parameters for the mercury atoms has also been observed for the red and orange HgI₂ structures determined from accurate single crystal data [6,7].

4. Conclusions

The diffraction experiments have shown the existence of an additional phase (V) of HgBr₂ above 9.0 GPa. Together with published material [8,9], they suggest a phase transition.
sequence, (I) orthorhombic–(II) orthorhombic–(III) monoclinic–(IV) trigonal–(V) trigonal or hexagonal. Due to strong texture effects, we cannot unambiguously confirm or disprove the existence of the I–II transformation, and we are unable to index the pattern and ascertain the proposed symmetry of phase III. The destructive character of the III–IV transition considerably reduces the texture effects, allowing a complete structure determination of phase IV. This phase forms a commensurately modulated CdI$_2$ structure built from layers of edge-sharing octahedra, in which one out of three mercury atoms assumes a trigonal rather than an octahedral coordination. This structure and its atomic displacement parameters suggest mercury to be mobile and recall the transformation of orange into red HgI$_2$ where mercury also appears to be mobile inside the iodine-atom substructure and diffuses between tetrahedral voids [6]. The structure of phase IV of HgBr$_2$ may explain the shift of the Hg–Br Raman frequencies to lower values with increasing pressure [12]. Usually, a vibrational mode shifts to higher frequency upon increasing pressure. The structural reason for the decreasing frequency observed for HgBr$_2$ may be the tendency of mercury to shift to higher coordination with increasing pressure.

The results presented here are a further confirmation that Hg(II) can adopt a considerable variety of environments in condensed matter, including tetrahedral, trigonal, linear and octahedral coordination. Each coordination may derive from a region of stability in the $p$–$T$ space associated with particular values of the energy levels of the valence-electrons. For HgBr$_2$ the low pressure region of the phase diagram seems to favor molecular structures with a coordination number of two, while pressures above 4.0 GPa stabilize pyramidal and octahedral coordination with trigonal and hexagonal symmetries.

The halides of the other group IIB elements Zn and Cd form non-molecular structures with closest-packed anions. The structures of the zinc halides are tetrahedral [20] analogous (but not identical) to those of HgI$_2$, whereas the cadmium halides form the well-known octahedral layer structures of which phase IV of HgBr$_2$ is a variant.

Acknowledgements

We thank Prof. Majed Chergui of the Laboratoire de spectroscopie ultrarapide, Swiss Federal Institute of Technology at Lausanne for providing the setup used in the SHG experiments. We also thank the staff of the Swiss–Norwegian Beam Lines at the ESRF, Grenoble, for their competent help. This work, as well as in part the Swiss–Norwegian Beam Lines, has been funded by the Swiss National Science Foundation.

References

GSAS profile function for constant wavelength X-ray data was used, and two overall isotropic atomic displacement parameters, one for mercury and one for bromine, were included in the refinement. Texture effects were taken into account using a March–Dollase texture model. Contamination by the gasket was modelled by incorporating tungsten as an additional phase.

[19] Details were as follows: the background was modeled using a Chebyshev polynomial function with 14 terms, the default