X-Ray Raman Spectroscopy of High-Pressure Phases and Novel Materials derived from Low-Z Compounds

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Introduction

We have used X-ray Raman spectroscopy (XRS) to study the oxygen and carbon K-edges of high pressure phases of CO and CO2 to understand changes in chemical bonding associated with the pressure-induced molecular-to-nonmolecular phase transitions in these materials. This work has exploited the power of the XRS technique to resolve important scientific questions regarding the nature of the high-pressure phases of CO and CO2.

The application of high pressures often transforms molecular solids to nonmolecular extended solids, manifesting entirely different chemical bondings and crystal and electronic band structures. Because of a sizeable difference in bondings and a large activational energy, these novel forms of high-pressure phases are often metastably recoverable, yet inaccessible to conventional chemical synthesis techniques at ambient pressures. These new phases are often of geophysical interest because the pressure/temperature-synthesis conditions are relevant to planetary interiors and in many cases the crystal structures are commonly shared by terrestrial minerals. The classic example of such a system is diamond, and recent work, including research of the proposing group, has found new examples including novel phases derived from Carbon Monoxide [1] and Carbon Dioxide [2]. Kinetic barriers at high pressures often hinder the transformations leading to incomplete or disordered products. Thus x-ray diffraction may not definitively identify the new phase and additional diagnostic studies can serve to validate phase and structural identifications. Recent x-ray Raman studies of the oxygen K edge by Bergmann et al. at APS Sector 18 demonstrate its power in studying the local structure of water molecules [3].

At pressures of ~50 kbar, the CO molecular solid transforms to a novel phase that loses its molecular signature and, we contend, interconnects to form a covalently bond network (extended solid) [1]. The high-pressure phase of CO can be quenched and is metastable at ambient pressure and temperature. The ambient condition metastability/lifetime of the CO-derived solid can vary from minutes to seemingly indefinite.

We hypothesize that this variation in lifetime is related to the degree of polymerization. We have performed various optical spectroscopies on this material, but a definitive conclusion remains elusive and the absorption band of the diamond anvil obscures important regions of the infrared spectrum. The recovered materials also appear to be amorphous, so x-ray diffraction provides only limited information. Using XRS we will study the oxygen and carbon K-edges to observe time-dependent changes at high pressure and in the quenched metastable phase. Information on the electronic environment of the oxygen and carbon atoms can yield insight into both the transformation and decomposition processes. This material is energetic and may serve as a high-density fuel for propulsion applications [4].

In our recent studies, we have discovered that molecular CO2 (cubic phase I) transforms to a covalently bonded extended solid (tridymite-like phase V) with four-fold coordinated carbon) above 40 GPa and 2000 K [2]. This transformation is believed to occur as a result of pressure-induced electron delocalization. However, the exact mechanism is not very well understood as yet. For example, the presence of intermediate phases such as pseudo-six-fold CO2-phase II and bent CO2-phase IV suggests that the electron delocalization apparently occurs continuously with increasing pressures above 10-20 GPa [5,6].

Fig. 2. Phase diagram of CO2. Blue lines established by experiment, red lines indicate theoretical predictions. Adapted from ref.s 6 and 7.

A recent theoretical work, on the other hand, suggests a quite different “abrupt” mechanism [7]. One way to clarify this discrepancy is to directly determine the change in chemical bonding in carbon dioxide at high pressures, which should be reflected in the near-K-edge absorption spectra of oxygen and carbon.

Both carbon and oxygen have very low energy x-ray absorption bands, ~290 eV and ~540 eV for the carbon and oxygen K-edges respectively, which present a severe challenge for x-ray absorption spectroscopy. The x-rays at this energy cannot
penetrate through the relatively thick and absorptive samples and diamond anvils. However, this limitation can be overcome using the XRS technique of measuring inelastically scattered high-energy x-rays. The performance characteristics of a high-intensity third-generation synchrotron source like the APS are ideally suited and crucial to this experiment. Several sectors at the APS, including Sectors 18 and 13, have demonstrated such experiments in several low-Z materials like H2O [3] and carbon [8] at both ambient and high pressures.

**Methods and Materials**

In this study, CO2 and CO samples were be loaded cryogenically in diamond anvil cells with beryllium gaskets (x-ray transparent) and a few ruby particles for sensing the pressure. Typical sample sizes were 30 µm x 100 µm (thickness, diameter). Diamond anvil cells manufactured with a large conical aperture perpendicular to the load axis were used. The x-ray beam (~100 µm spot size) was incident upon the sample through the a small hole in the DAC and then the beryllium gasket. The scattered x-rays were collected at ~30 degrees from the incident beam. The conical aperture permitted collection of a large solid angle of x-rays scattering and exiting through the gasket. The incident high-energy monochromatic x-ray was at an energy of ~10 keV. Following the work of previous researchers [3,9], we will scan the incident x-ray monochromator over a range of 50-100 eV. The inelastically scattered photons were detected using a six-crystal analyzer (resoln. 1.7eV) at a fixed energy, E0, using a multi-crystal analyzer and a liquid-nitrogen-cooled Ge detector. Thus the Raman shift \( \Delta E = E_f - E_0 \) is determined by scanning \( E_0 \) rather than the conventional optical approach of a fixed \( E_0 \) and a scanned \( E_f \). Typical times for collection of XRS spectra were 6-8 hours.

**Results**

Oxygen and Carbon K-edge XRS spectra were successfully collected for CO2. Unfortunately the 10 keV x-ray induced radiation damage and/or the transformation of CO to an unidentified phase, possibly related to a polymeric phase of CO that has been produced at high pressure [4]. Based on an independent set of experiments, we believe this transformation occurs quite rapidly, on timescales of minutes to hours. Because of the long collection times the XRS spectra of the CO is certainly a mixture of pure CO and transformed material with unknown relative strengths.

Carbon k-edge XRS spectra are to be treated with some reservations since the carbon in the diamond anvil cannot be eliminated and may enter into the measured data.

**Discussion**

The XRS oxygen k-edge spectra are the most readily tractable data from our studies. These data do not have any radiation damage/conversion or diamond anvil contaminations issues.

Initial review of the XRS data identify clear differences in the spectra in the experimentally identified phases, supporting the contention of changes in not only crystal structure but chemical bonding in phases II, III, and IV (associated molecular vs molecular). Clear differences are observed in the spectra, and detailed analysis should yield information on the local structure of these phases. Continued study and analysis is underway including

- improved studies (larger samples, higher S/N) of CO2 (II, IV &V), particularly the polymeric phase V.
- develop a more quantitative and theory-based interpretation of XRS spectra in relation to electronic changes in high-pressure phases.
- study pressure dependence of XRS within phases as compared to across phase lines.
• apply XRS to other molecular systems of interest.

Acknowledgements
This work was performed under the auspices of the U.S. DOE by the Univ. of Calif., LLNL under contract No. W-7405-Eng-48 and at GeoSoilEnviroCARS, by the NSF- Earth Sciences (EAR-0217473), DOE - Geosciences (DE-FG02-94ER14466) and the State of Ill. Use of the Advanced Photon Source was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract No. W-31-109-ENG-38.

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