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ON THE ELASTIC BEHAVIOUR OF ZEOLITE PAULINGITE: A SYNCHROTRON POWDER DIFFRACTION STUDY

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The high-pressure (HP) elastic behaviour of a zeolite paulingite found in vugs in olivine nephelinite of the Vinařická hora Hill Tertiary volcano near Kladno (Czech Republic) was studied by means of in situ synchrotron X-ray powder diffraction. The diffraction experiments were performed using a diamond anvil cell (DAC) at the X7A beamline at the National Synchrotron Light Source at Brookhaven National Laboratory. A gas-proportional position-sensitive detector was stepped in 0.25° intervals over the angular range of $3\text{--}35^\circ$ with counting times of 90 - 150 s per step. The wavelength of the incident beam (0.60046 \AA), PSD zero channel, and PSD degrees per channel were determined from a CeO_2 standard (SRM 674). A powdered sample of paulingite was placed in the gasket hole of the DAC together with some ruby chips. The pressure at the sample was measured by detecting the shift in the R1 emission line of the ruby chips with precision $\pm 0.1 \text{ GPa}$. Two sets of experiments were carried out with two different pressure-transmitting media: a nominally penetrating methanol : ethanol : water (16:3:1) mixture and a nominally

non-penetrating silicon oil. The maximum pressures attained in the experiments were 5.12 GPa in the case of alcohol : water mixture and 2.53 GPa in the case of silicon oil. The chemical composition of the studied sample, $\text{K}_{4.87}\text{Na}_{0.37}\text{Ca}_{3.20}\text{Ba}_{0.23}\text{Mg}_{0.04}\text{Sr}_{0.03}\text{Fe}_{0.05}\text{Al}_{11.17}\text{Si}_{30.52}\text{O}_{84} \cdot 30 \text{ H}_2\text{O}$, has been determined by an electron microprobe analyser CAMECA SX-100. Unit cell parameters were determined by LeBail whole powder pattern fitting implemented in the FullProf suite. Axial and volume bulk moduli have been calculated using a truncated second-order Birch–Murnaghan equation-of-state using program EOSFIT. No phase transition or sample decomposition have been observed up to the maximum recorded pressures. The bulk moduli refined from the data collected in nominally penetrating medium were excessively high pointing out that paulingite is over-hydrated under the conditions of the experiment. The refined elastic parameters for the experiment with nominally non-penetrating medium are: $V_0 = 43794(185) \text{ \AA}^3$, $K_{T0} = 18(1) \text{ GPa}$.

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SYNTHESIS AND CRYSTAL STRUCTURES OF $\text{Pd}_3\text{Pb}_2\text{Te}_2$ AND Pd_3AgSe

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This presentation is a part of systematic research on crystal structures, phase relations and selected physical properties of phases from Pd-Pb-Te and Pd-Ag-Se ternary systems. These systems comprise many platinum-group minerals or phases, which can be found in various ore deposits as potential new minerals. Consequently, phases from these systems are of interest in mineralogy and geology of ore deposits.

The ternary compounds $\text{Pd}_3\text{Pb}_2\text{Te}_2$ and Pd_3AgSe were synthesised from elements by conventional solid-state reactions. Stoichiometric amounts of individual elements were sealed in evacuated silica glass tubes and resultant mixtures were heated at selected temperatures. Samples with $\text{Pd}_3\text{Pb}_2\text{Te}_2$ and Pd_3AgSe were heated at 400°C and 350°C , respectively. After long-term annealing, the samples were quenched in a cold-water bath. All attempts to

prepare single crystals of studied phases suitable for single-crystal analysis failed, the crystal structure determinations of $\text{Pd}_3\text{Pb}_2\text{Te}_2$ and Pd_3AgSe were performed from conventional powder X-ray diffraction data. The structure solutions of title compounds by direct methods were accomplished using the program EXPO2004 [1]; subsequent refinements were performed with FullProf program.

$\text{Pd}_3\text{Pb}_2\text{Te}_2$: Space group $Pmmn$, $a = 8.59$, $b = 5.93$, $c = 6.31 \text{ \AA}$, $V = 322 \text{ \AA}^3$ and $Z = 2$. Its crystal structure can be described as a layered structure formed by face-shared $[\text{PdPb}_4\text{Te}_2]$ octahedra running parallel to \mathbf{b} axis. Two independent palladium atoms are surrounded by four lead and two tellurium atoms showing distorted octahedral coordination with tellurium atoms in *trans* positions with respect to one another. Two independent lead atoms are coordinated by six palladium atom in two different ways. The