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**Determination of Equations of State for AlF<sub>3</sub> and AlI<sub>3</sub>: Semi-empirical Modeling of Extreme Condition Halide Chemistry** JOSEPH ZAUG, Lawrence Livermore National Laboratory, ELISSAIOS STAVROU, Carnegie Institute of Washington, Geophysical Laboratory, SORIN BASTEIA, JONATHAN CROWHURST, Lawrence Livermore National Laboratory, ALEXANDER GONCHAROV, Carnegie Institute of Washington, Geophysical Laboratory, SARAH ROBERTS, JONATHAN PLAUE, JEFFREY CARTER, MICHAEL ARMSTRONG, Lawrence Livermore National Laboratory — Pressure dependent angle-dispersive x-ray powder diffraction measurements of alpha-phase aluminum trifluoride (alpha-AlF<sub>3</sub>) and separately, aluminum triiodide (AlI<sub>3</sub>) were conducted using a diamond-anvil cell. Results at 295 K extend to 50 GPa. The equations of state of AlF<sub>3</sub> and AlI<sub>3</sub> were determined through refinements of collected x-ray patterns. The respective bulk moduli and corresponding pressure derivatives using multiple orders of the Birch-Murnaghan, Ff, and Gg EoS models will be discussed. Aluminum trifluoride exhibits no pressure induced structural phase transition while the triiodide data reveal a second-order iso-structural rearrangement: Applied stress transformed a monoclinically distorted face centered cubic (FCC) structure into a perfect FCC structure. Results from semi-empirical thermochemical computations of energetic materials formulated with fluorine containing reactants will be presented. \* This work was performed under the auspices of the U.S. Department of Energy jointly by Lawrence Livermore National Laboratory under Contract DE-AC52-07NA27344.

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