DEVELOPING V-XANES OXYBAROMETRY FOR PROBING MATERIALS FORMED IN REDUCING ENVIRONMENTS IN THE EARLY SOLAR DISK. A.L. Butterworth¹, Z. Gainsforth¹, C. E. Jilly-Rehak¹, K. Righter², and A. J. Westphal¹. ¹University of California Berkeley, Space Sciences Laboratory, 7 Gauss Way, Berkeley, CA 94720. ²Johnson Space Center, Houston, TX 77058. Email: <u>annab@ssl.berkeley.edu</u>.

Introduction: Vanadium exhibits four oxidation states (V^{2+} , V^{3+} , V^{4+} , and V^{5+}) that have been shown to preferentially partition between melt phases dependent on redox conditions, spanning oxygen fugacity across more than 10 log units [1]. We are developing synchrotron-based x-ray absorption spectroscopy of low-fugacity standards for the determination of V oxidation state in highly reducing conditions relevant to the early solar nebula.

Development of V oxybarometers has progressed significantly in recent years, particularly for V³⁺, V⁴⁺ and V⁵⁺, equivalent to environments with log(fO₂) > -12. A V K-edge XANES oxybarometer was calibrated (@1400°C) with glasses [2] then expanded to chromian spinels [3] and spinel-olivine-glass melts [4].

K-edge based oxybarometers depend on preedge features (1s–3d transitions) and edge position to probe oxidation states. Advantages of a Kedge based oxybarometer includes the relative simplicity of interpreting pre-edge features. Modeling tools to interpret K-edge spectra have been well established, for example first-principles calculations based on the density functional theory (DFT) [5].

One limitation of the V K-edge oxybarometer is the calibrated oxygen fugacity range. The glasses available to constrain the initial calibrations range between log (fO₂) ~ -5 (mainly V⁵⁺) down to log (fO₂) ~ -12, at 1400°C, (mainly V³⁺). The oxybarometer [2] extrapolated from V³⁺ to V²⁺ assuming a V²⁺ pre-edge peak height of zero.

Another potential limitation of K-edge synchrotron techniques is spatial resolution. High resolution, sub-micron hard X-ray *microXANES* applications (approaching 100 nm beamspot) must still consider several- μ m penetration depth. Thicker rock sections or sample fine-scale heterogeneity may hide sub-surface inclusions that could contaminate spectra.

L-edge techniques include soft X-ray synchrotron XANES and TEM EELS. Both techniques are capable of higher spatial resolution than K-edge XANES (<10 nm) at V L-edge. The $L_{2,3}$ 2p-3d multiplet structure is sharp and more sensitive to combined effects of valence state, site symmetry, and crystal field strength than K-edge features. V²⁺ L-edge spectra also show strong features in contrast to 1s-3d preedge. This is especially important for low abundance V (<0.3 wt%) in primitive meteorite spinels.

Interpretation of L-edge spectroscopy is generally less well-developed than for K-edge simulations. In contrast with the K-edge approach, the entire L₂₃ spectrum must be analyzed for interpretation. High quality ab initio calculations of the Ledge of metal oxide compounds, including Ti and V, are being actively developed [6,7]. So far, models apply to a relatively narrow range of mineral structures and compositions, meaning no universal models exist for V L_{2.3} spectra. As far as we know, there is no tool to determine precise (fO₂ determination within $\pm 0.5 \log \text{ units}$) \tilde{V}^{2+}/V^{3+} calibrated for spinels appropriate for probing micro-environments of primitive, reduced solar system materials (such as spinels in chondrules and CAIs).

Our approach is to continue the development of V, Ti oxybarometers for astromaterials, focusing on magnesian spinel formed in reducing environments, log (fO_2) < -12. We are beginning with existing standards [3], and also synthesizing new standards, with V²⁺ or Ti³⁺, to probe lower fO₂.

Samples and Methods: We have analyzed chromian spinel samples, which were previously used to extend the V K-edge oxybarometer [3], Table 1. We acquired multi element spectra at both K- and L-edges: Fe, Cr, Ti, V K-edge XANES and EXAFS at ALS Beamline 10.3.2, and Fe, Cr, Ti, V L-edges plus Al, O K-edges XANES at ALS 11.0.2. For μ XAS Beamline 10.3.2, we used the original polished samples of spinel/melt pairs embedded in epoxy. We prepared FIB sections of these samples for L-edge work at ALS 11.0.2 (Scanning Transmission Xray Microscope, STXM) and TEM. FIB and TEM work was carried out at the National Center for Electron. FIB liftouts were cut using an FEI DualBeam FIB with Ga⁺ at 30 and 6 keV. Atom Location by CHanelling Enhanced Microanalysis (ALCHEMI) was done on the Titan to determine the sites of cations.

Using this approach we aim to anchor our V K-spectra to previous results, and enabling quantitative interpretaion of L-edge spectra.

Spinel/Melt Buffer & Dopant	Log fO ₂ , 1300°C	ΔIW 1300°C
IW	-11.15	-0.40
IW, V+5%	-11.15	-0.40
IW, Ti+10%	-11.13	-0.38
IW-1	-12.09	-1.51

Table 1: Samples analyzed, from [3]

 $\overline{IW} = Iron-W$ üstite buffer

The IW-1 sample microprobe characterization was (in wt%) 43.3 Cr_2O_3 , 1.0 V_2O_3 . 16.22 FeO, 13.99 MgO [3].

Results: Figs 1 and 2 show example V spectra taken from one sample, "Experiment IW-1", which is our lowest fO2 available standard. In Fig. 1, we also show for comparison an L-edge XANES spectrum for Stardust particle "Iris", thought to be a fragment of a Type II chondrule [8]. The spectra are qualitatively similar, predominantly V^{3+} .

We now have a complete description of the four chromian spinel-structure oxides, including formation fO_2 , composition, site occupancy and multi-element K-edge spectra, we will use DFT and multiplet theory to simulate each of the spinels. Vanadium composition ranges from major element to minor abundance.

Discussion: The K-edge simulations will provide the basis for developing a model for L-edge interpretation. We are preparing low oxygen fugacity spinel standards at JSC, aiming to create samples with significant V^{2+} (and Ti³⁺ [9]). Subsequent availability of lower fO₂ V-bearing spinel samples will then allow us to extend the range of the K-edge oxybarometer.

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References: [1] Papike, P. V. et al., (2005) *Am. Min.* 90, 277.. [2] Sutton S. R. et al. (2005) *GCA* 69, 2333-2348. [3] Righter K. et al. (2016) *Am. Min. 101*, 1928-1942. [4] Papike, P. V. et al., (2013) *Am. Min.* 98, 2193. [5] Cabaret, D. et al., (2010) *Phys. Chem. Chem. Phys.*, 12, 5619. [6] Hoche T., et al., (2013) Am. Min. 98, 665. [7] Brik M. G., et al., (2006). *The Euro. Phys. J. B* 51: 345. [8] Gainsforth Z. et al, (2015), *MaPS* 50, 976. [9] Righter et al., (2017) LPSC.

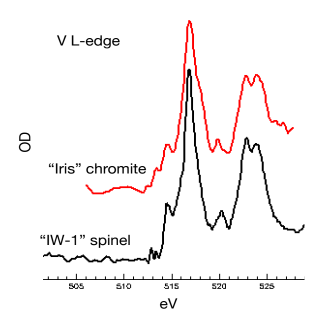


Fig 1. V L- XANES spectra of Stardust particle "Iris" [8] and qualitatively similar spectra acquired on chromian spinel "IW-1" from [3].

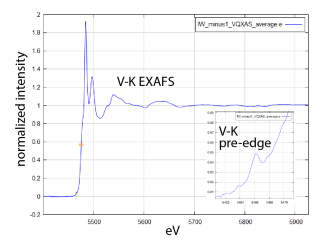


Fig 2. V K-egde spectra the same chromian spinel "IW-1" from [3]. Spectrum is high resolution EXAFS, the inset shows pre-edge XANES region for comparison with [1]).