

**L-EDGE XANES MEASUREMENTS OF THE OXIDATION STATE OF TUNGSTEN IN IRON BEARING AND IRON FREE SILICATE GLASSES.** L. R. Danielson<sup>1,2</sup>, K. Righter<sup>2</sup>, S. Sutton<sup>3</sup>, M. Newville<sup>3</sup>, L. Le<sup>1,2</sup>, <sup>1</sup>Jacobs Sverdrup Co., Houston, TX 77058 United States (lisa.r.danielson@nasa.gov), <sup>2</sup>NASA JSC, 2101 NASA Road One, Houston, TX 77058 United States, <sup>3</sup>GSECARS University of Chicago, 9700 South Cass Avenue, Bldg. 434A, Argonne, IL 60439 United States.

**Introduction:** Tungsten is important in constraining core formation of the Earth because this element is a moderately siderophile element (depleted ~ 10 relative to chondrites) and, as a member of the Hf-W isotopic system, it is useful in constraining the timing of core formation. A number of previous experimental studies have been carried out to determine the silicate solubility and metal-silicate partitioning behavior of W, including its concomitant oxidation state. However, results of previous studies (Fig. 1) are inconsistent on whether W occurs as  $W^{4+}$  or  $W^{6+}$ .

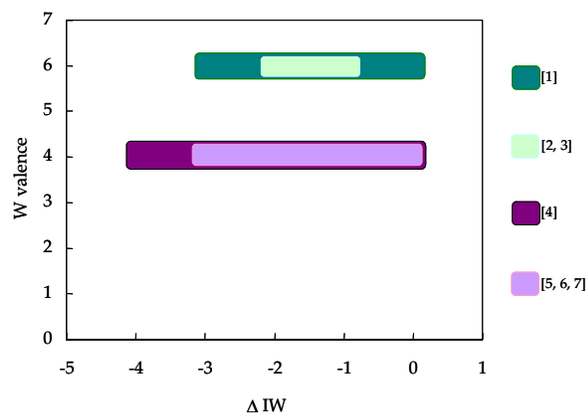


Figure 1. Comparison of W valence from previous results.

It is assumed that  $W^{4+}$  is the cation valence relevant to core formation [8]. Given the sensitivity to silicate composition of high valence cations [8], knowledge of the oxidation state of W over a wide range of  $fO_2$  is critical to understanding the oxidation state of the mantle and core formation processes. This study seeks to measure the W valence and change in valence state over the range of  $fO_2$  most relevant to core formation, around IW-2.

**Experiments:** Two compositions were used to determine the effects of iron content. Initial experiments, reported in [9] were conducted at 1300 °C, for durations of 24 to 96 hours and air or water quenched. One series was conducted using the An-Di eutectic, from QFM to IW-5 ( $\log fO_2$  -7.25 to -16). Experiments using an ankaramite starting composition were conducted from IW-1 to IW-5 ( $\log fO_2$  -11.75 to -16). Experiments were doped with 1wt% of  $WO_3$ . For both starting compositions, at IW-1, one set of experiments was doped with 1wt% of  $WO_2$ . Experiments at IW-2 and

above were conducted using a Re wire loop technique, while experiments below IW-2 were conducted in sealed silica tubes as in Fig. 2.

The most recent experiments reported herein expanded the  $fO_2$  range, from in air to IW-10 (ankaramite runs up to  $\log fO_2$  -0.4, CMAS runs down to  $\log fO_2$  -20) of the data set, and were designed to in an attempt to improve success with analyzable runs at low  $fO_2$ . Run duration was increased to 120 hours and all experiments were water quenched. Because the solubility of W decreases to ~100 ppmw [4] at the low  $fO_2$  conditions of these experiments, amount of W as  $WO_3$  added was below 1000 ppmw.

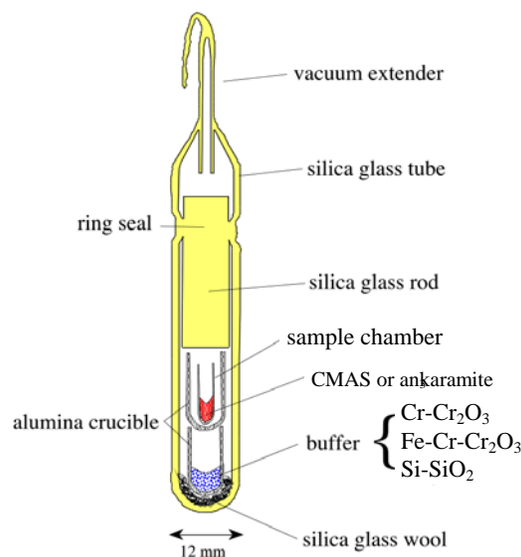


Figure 2. Schematic of sealed silica tube experiments conducted at lowest  $fO_2$ , below IW-2.

**Analytical:** A monochromatic X-ray beam from a Si(111) double crystal monochromator was focused onto the sample and the fluorescent X-ray yield was plotted as a function of incident X-ray energy (more detail can be found in [10]). The oxidation state of tungsten was inferred from the energy of the first peak in the LIII-edge derivative spectrum.  $WO_2$ ,  $WO_3$ ,  $FeWO_4$ ,  $CaWO_4$ , were used as standards.

**Results:** Results (Fig. 3 and 4) for both the iron-bearing and iron-free starting materials suggest that only  $W^{6+}$  is present from the most oxidized conditions to IW ( $\log fO_2$  -10.75). At IW, tungsten starts to exhibit mixed valence but is still dominated by  $W^{6+}$  for

ankaramite. At IW-2,  $W^{4+}$  becomes more abundant for ankaramite, but a definitive transition to  $W^{4+}$  below IW-2 has not been observed. CMAS has a lower W valence than ankaramite below IW, with the most reduced state observed being  $W^{4+}$ . These results suggest that  $W^{4+}$  would be dominant below IW-2.

Even with improved experimental technique focused on reducing the “nugget effect,” at IW-2 and below, metal exsolved from the silicate, complicating the analyses. Analyses in nugget free regions of the silicate portion of the experiments cluster around  $W^{5+}$ , while analyses in nugget-rich regions cluster around  $W^{2+}$ .

Four CMAS run products from Ertel [4] were also analyzed and shown for comparison in Fig. 4. These samples were reported as  $W^{4+}$ , from the slope of the W concentration vs.  $\log fO_2$  line. The Ertel run products are in good agreement with our CMAS run products, showing decreasing W valence with decreasing  $fO_2$ , but appear to be mixed  $W^{6+}$  and  $W^{4+}$  valence.

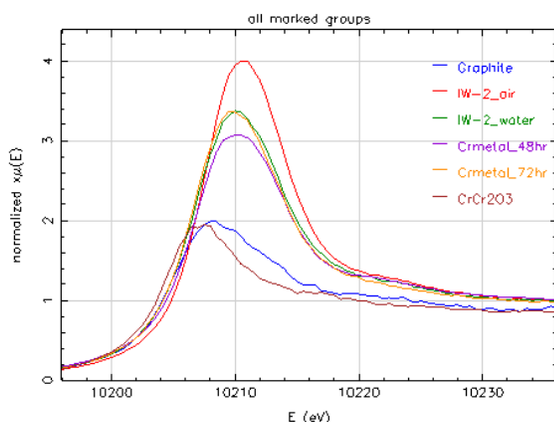


Figure 3. A sampling of results from some CMAS run product analyses (figure 4), showing range of W valence, from  $W^{6+}$  (IW+1 air quench, red line) to around  $W^{2+}$  (IW-5, using Cr-Cr<sub>2</sub>O<sub>3</sub> buffer, brick line). Data show differences between air (red line) and water quench (green line) at IW-2, with a 0.5+ reduction in W valence observed for a water quench. Other lines indicate a run in a graphite capsule, and runs using only Cr metal as a buffer.

**Discussion and Conclusions:** Both CMAS and ankaramite glasses show  $W^{6+}$  above IW and mixed valence below IW. The transition to  $W^{4+}$  only, appear to happen at or below IW-2 for iron free systems, but probably below IW-3 for iron bearing systems. Analyses of run products from previous experiments [4], support the observation the iron free transition to only  $W^{4+}$  occurs at or below IW-2.

The mixed states below IW-2 may result from analyses in which both silicate glass and exsolved W-bearing metal are present in the analytical volume in

varying proportions. This suggests if all W were dissolved in the silicate in a nugget free condition, W valence would be less than  $W^{5+}$  at lowest  $fO_2$ .

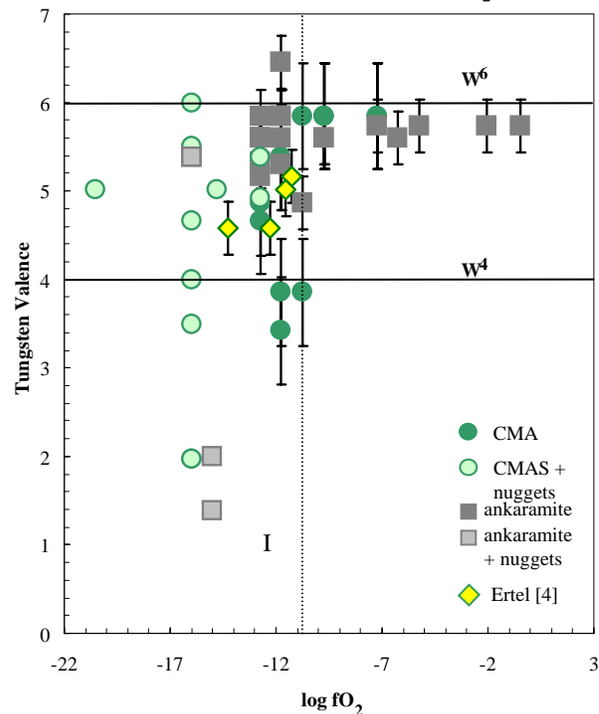


Figure 4. Summary of valence results inferred from the energy of the first peak in the LIII-edge derivative spectrum.

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