**EVIDENCE FOR INTERLAYER COLLAPSE OF NONTRONITE ON MARS FROM LABORATORY VISIBLE AND NEAR-IR REFLECTANCE SPECTRA.** R. V. Morris¹, D. W. Ming², D. C. Golden², T. G. Graff², and C. N. Achilles², ¹ARES NASA Johnson Space Center, Houston, TX 77058, richard.v.morris@nasa.gov and ²Jacobs Engineering, ESCG, Houston, TX.

**Introduction:** Dioctahedral smectites (e.g., nontronite and montmorillonite) are interpreted to occupy the optical surface of Mars at a number of locations on the basis of spectral features derived from interlayer H₂O and MOH (M=Fe³⁺, Fe⁵⁺Al, Al₂, etc.) as observed by orbiting MRO-CRISM and MEx-OMEGA hyperspectral imaging spectrometers [e.g., 1-4]. At wavelengths shorter than ~2.7 µm, the strongest bands from interlayer H₂O occur at ~1.4 and 1.9 µm from 2v₁ and v₁+v₂, respectively, where v₁ and v₂ are the fundamental stretching and bending vibrations of the H₂O molecule [5]. Smectite MOH vibrations occur near 1.4 µm (stretching overtone) and in the region between 2.1 and 2.7 µm (stretching + bending combination). Because interlayer H₂O can exchange with the martian environment [e.g., 6], a number of studies have examined the strength of the interlayer H₂O spectral features under Mars-like environmental conditions [7-9]. The relationship between spectral properties and the underlying crystal structure of the smectites was not determined, and the extent of interlayer H₂O removal was not established.

We report combined visible and near-IR (VNIR), Mössbauer (MB), and powder X-ray diffraction (XRD) data for samples of the Fe-bearing smectite nontronite where the interlayer was collapsed by complete removal of interlayer H₂O.

**Samples and Methods:** Pennsylvania nontronite (PHY07, <38 µm [10], courtesy of T. Roush) and the Garfield Washington nontronite (API33A, <150 µm) with their native exchangeable cations were used for our experiments.

VNIR reflectivity spectra (0.35-2.50 µm) were acquired with an Analytical Spectral Devices (ASD) FieldSpec3 fiber optic spectrometer configured with a Mug light. The instrument was co-located with a hot plate, dewpoint meter (Vaisala DRYCAP DM70), and an IR thermometer (Fluke Model 66) in a 1-atm. glove box. The well-defined Fe³⁺ electronic bands at ~0.45, 0.63, and 0.95 µm for the unheated nontronites broaden and increase in intensity (surface darkens) with increasing temperature and time at low environmental H₂O concentration, the intensity of the 1.4 and 1.9 µm bands from interlayer H₂O decrease (in agreement with [1-4]) and, after heating at 210°C, are virtually not detectable. The spectral features at 2.29 µm and 2.40 µm associated with Fe³⁺-OH do not change position and are better resolved as interlayer H₂O is lost. The band at 1.44 µm is the overtone for the Fe³⁺-OH stretching vibration. XRD patterns were obtained in ambient air within 7 min after their removal from the glove box.

**Results:** The XRD patterns for API33A and PHY07 (Figs. 1 and 2) show that heating at ~210°C and a final concentration of 100-130 ppmv H₂O in the glove box resulted in full interlayer collapse (blue traces) as indicated by the (001) basal diffraction peak at 9.5 Å [e.g., 11]. After ~2 hr in air (red traces), the nontronites acquired interlayer H₂O as indicated by the basal diffraction peak at ~14.6 Å, although the basal peaks are broad compared to unheated samples (green traces). The basic smectite crystal structure has not been altered, however, because non-basal diffraction peaks (020) and (060) at 4.51 Å and 1.52 Å [11], respectively, are essentially invariant with respect to position and width.

The MB spectrum for PHY07 heated to 210°C shows no detectable Fe⁷⁺ (Fig. 3). The spectrum is well fit by two octahedral Fe⁷⁺ doublets (from [e.g., 12]) and no tetrahedral Fe⁵⁺ doublet.

The nontronite VNIR spectra are dramatically effected by interlayer collapse (Figs. 4 and 5). With increasing temperature and time at low environmental H₂O concentration, the intensity of the 1.4 and 1.9 µm bands from interlayer H₂O decrease (in agreement with [1-4]) and, after heating at 210°C, are virtually not detectable. The spectral features at 2.29 µm and 2.40 µm associated with Fe³⁺-OH do not change position and are better resolved as interlayer H₂O is lost. The band at 1.44 µm is the overtone for the Fe³⁺-OH stretching vibration.

The well-defined Fe³⁺ electronic bands at ~0.45, 0.63, and 0.95 µm for the unheated nontronites broaden and increase in intensity (surface darkens) with increasing time/temperature until, with complete interlayer collapse, a positive, featureless slope is present between ~0.5 and 2.2 µm. We interpret progressive darkening of the surface in part to progressive removal of interlayer H₂O and in part to the transition of the dark layer from optically thin to optically thick. Upon exposure of the collapsed nontronites to lab air, interlayer H₂O is acquired as evidenced by the 1.4 and 1.9 µm bands, but the Fe³⁺ spectral features still closely resemble those for the collapsed nontronites even after ~744 hr of exposure to air for PHY07 after heating in dry N₂ at 210°C (Fig. 6).
Application to CRISM, OMEGA, and MSL-CheMin: Our spectra for collapsed nontronite have virtually no spectral features for interlayer H₂O at ~1.4 and 1.9 μm, retain the spectral features from Fe³⁺OH with enhanced spectral contrast at the same locations (2.29 and 2.41 μm), and have a positive, low-albedo spectral slope instead of well-defined bands from Fe³⁺ between ~0.5 and 2.0 μm. Thus, the low relative intensity of H₂O bands compared to Fe³⁺OH bands and the absence of well-defined Fe³⁺ bands (e.g., at 0.96 μm) for martian nontronite are evidence for significant loss of interlayer H₂O if not interlayer collapse in response to arid environmental conditions. The detection of a 9.5-10 Å diffraction peak by MSL-CheMin might be in situ evidence for collapsed nontronite.