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# Spectral reflectance properties of zeolites and remote sensing implications

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[1] The 0.3- to 26- $\mu$ m reflectance spectra of a suite of 28 zeolites were measured and analyzed to derive spectral-compositional-structural relationships. Below  $\sim 7 \,\mu m$ , the spectra are largely dominated by absorption features associated with zeolitic water. At longer wavelengths, the spectra are dominated by absorption features associated with the aluminosilicate framework. The spectra exhibit a number of systematic variations which can be used for both structural and compositional determinations. These include: (1) distinguishing different structural groups on the basis of wavelength position variations associated with absorption features in the 8.5- to  $26-\mu m$  region that are related to differences in the structure of the aluminosilicate framework; (2) determining the major cation which is present (Ca, Na, K) and the associated electronic environment of the zeolitic water on the basis of how these cations hydrogen bond to the water molecules in the void spaces and consequently affect water-related absorption band positions, particularly in the 1.4, 1.9, and 2.0- to 2.5- $\mu$ m regions; (3) determining the Al:(Al + Si) ratio and SCFM chemical index on the basis of absorption features in the 7- to 26-um region which are most sensitive to these compositional variations; and (4) identifying ironbearing zeolites on the basis of absorption features in the 0.35- to 0.9-µm region. The wavelength position and number of H<sub>2</sub>O-associated absorption bands are sensitive to factors such as the type of major cation, degree of hydrogen bonding, and size of the void space, all of which are somewhat interrelated. INDEX TERMS: 3934 Mineral Physics: Optical, infrared, and Raman spectroscopy; 5470 Planetology: Solid Surface Planets: Surface materials and properties; 6225 Planetology: Solar System Objects: Mars

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#### 1. Introduction

[2] Zeolites are an important class of minerals for a wide variety of applications. These include industrial uses such as molecular sieves for ion filtration and separation of a diversity of compounds, as an additive in paper and rubber, and as catalysts for numerous chemical processes [e.g., *Feins and Mullen*, 1970; *Breck*, 1974; *Sand and Mumpton*, 1978; *Tsitsishvili et al.*, 1992]. Zeolites are also petrogenetically important because many of them form under restricted ranges of temperature and/or pressure [e.g., *Coombs et al.*, 1959; *Zen*, 1961; *Breck*, 1974; *Sand and Mumpton*, 1978; *Barrer*, 1982; *Donahoe and Liou*, 1985; *Bowers and Burns*, 1990], and hence are potentially useful indicators of petrogenesis.

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[3] Zeolites have been suggested by a number of investigators as being plausibly present on the surface of Mars. There are a number of lines of indirect evidence supporting this contention, including their widespread presence in terrestrial basaltic and volcanic regions [Barnes et al., 1984; Browne et al., 1989; Zamarreno et al., 1989; Basu et al., 1998]. Zeolites also form in cold polar desert environments and are widely present in terrestrial cold desert soils, including those derived from basalts [Berkley, 1981; Berkley and Drake, 1981; Gibson et al., 1983; Gibson, 1985], as well as in the terrestrially weathered portions of Antarctic Martian meteorites [Gooding, 1984a, 1984b]. Zeolites may also form through low-temperature gradual alteration (palagonitization) of volcanic ash and tephra [Bonatti, 1965; Hay and Iijima, 1968; Geptner, 1970; Staudigel and Hart, 1983; Zhou and Fyfe, 1989; Jercinovic et al., 1990], and such a material is assumed to be widely present on the Martian surface [McSween and Murchie, 1999; Johnson et al., 1999; McSween et al., 1999; Bell et al., 2000].

[4] While zeolites can form in sedimentary environments [e.g., *Johnsson*, 1990] and possibly, on rare occasions, as primary igneous products [*Luhr and Kyser*, 1989], they are most commonly associated with hydrothermal environments [*Alt et al.*, 1986; *Iijima et al.*, 1984; *Foord et al.*,

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1986], including hydrothermally altered impact melt rocks and breccias [e.g., *Allen et al.*, 1982; *Golden et al.*, 1992; *Beiersdorfer and Ming*, 1996]. This is significant because impact cratering and hydrothermal alteration are, respectively, important known and inferred surface modification processes on Mars [e.g., *Carr*, 1996]. Zeolites can apparently form very quickly under appropriate hydrothermal conditions [*Browne et al.*, 1989].

[5] Other, less direct lines of evidence also suggest that zeolites may be present on the Martian surface. These include models of Martian weathering history and geological evolution [e.g., *Basu et al.*, 1998], and the behavior of the Martian atmosphere [e.g., *Gooding*, 1986]. As zeolites are able to incorporate large amounts of volatiles, such as  $CO_2$  and  $H_2O$  into their structure, they could be an important sink for a variety of Martian volatiles [*Gooding*, 1985; *Ming and Gooding*, 1988; *Jakes and Rajmon*, 1998]. Hence, detection of zeolites on Mars and determination of volatile contents and species would have important implications for reconstructing the planet's geological history, the evolution of its atmosphere, and fate of its inventory of volatiles [e.g., *Carr*, 1996].

[6] The current study focused on Na-, Ca-, and K-bearing zeolites. The 0.3- to 26-μm spectral reflectance properties of 28 zeolites, comprising 15 species, were examined in conjunction with compositional and structural information to determine the coupling between spectral variations, composition, and structure of this class of minerals. The goal was to determine which compositional and structural properties might be amenable to spectral analysis, allowing individual zeolites and zeolite structural and compositional groups to be spectrally distinguished from each other and from other minerals.

## 2. Structure, Composition, and Petrogenetic Significance of Zeolites

[7] The zeolite class of minerals includes a number of naturally occurring and synthetic species. Zeolites are characterized by an aluminosilicate three-dimensional framework [e.g., Deer et al., 1963] with different spatial arrangements of SiO<sub>4</sub> and AlO<sub>4</sub> tetrahedra. Individual zeolite minerals are distinguished on the basis of this spatial arrangement as well as their major cations and actual or potential volatile contents. We have adopted the classification of Breck [1974], who has subdivided zeolites into seven groups based on the specific array of (Al, Si)O<sub>4</sub> tetrahedra. Of these seven groups, six are represented by naturally occurring zeolites (group 3 contains only synthetic members). This structural classification was selected for the present study because of the close correspondence between structure and spectral properties for many minerals; it also accounts for other important properties such as void volumes and spatial arrangement of void spaces, cation exchange properties, and adsorption characteristics. At least two representatives from each of these six groups were included in this study.

[8] Zeolites are important minerals from the perspective of petrogenesis because many form under restricted conditions of pressure and/or temperature, and they have long been used to constrain petrogenetic conditions. The commonly cited "zeolite facies" is one manifestation of their petrogenetic usefulness [e.g., *Coombs et al.*, 1959; *Zen*, 1961]. Some of the formation conditions for the specific zeolites included in this study are summarized below (by structural groups as detailed by *Breck* [1974]). A more detailed discussion of zeolite formation conditions is given by *Barrer* [1982].

[9] The group 1 zeolites which were included in this study (wairakite, gismondine, laumontite, and phillipsite) are composed of single 4-ring (Al, Si)<sub>4</sub>O<sub>8</sub> units in various configurations (referred to as secondary building units). Differences between group 1 zeolites arise from the type of major cation present and the arrangement of the secondary building units [*Breck*, 1974]. The structural properties of these minerals are somewhat diverse; for example, laumontite contains both 4-, 6-, and 10-ring secondary building units. This group is also diverse in terms of the sites in which water molecules reside and the stability of the structure when subjected to dehydration [*Breck*, 1974].

[10] Wairakite generally forms at temperatures ranging between  $60^{\circ}$  and  $300^{\circ}$ C and at depths from 72 to 1600 m. Most wairakite forms at temperatures above 175°C (higher than most zeolites) although it has been reported to crystallize at surface temperatures [Kristmannsdottir and Tomasson, 1978]. Data from geothermal wells that pass through olivine basalt in Iceland indicate that gismondine crystallizes between 80° and 90°C [Kristmannsdottir and Tomasson, 1978]. The geobarometric conditions of formation for this mineral are not known. Laumontite has been considered an indicator mineral for high-temperature, depth zonation of zeolites in burial diagenesis, although it has also been found to form at very low temperatures [McCulloh et al., 1981]. Laumontite is very difficult to synthesize in the laboratory, requiring temperatures of 250°C and 1000 bars pressure [McCulloh et al., 1981]. Data from geothermal wells in basalt in Iceland indicate that laumontite crystallizes at temperatures between 98° and 230°C [Kristmannsdottir and Tomasson, 1978]. Laumontite is encountered in rhyolite, at temperatures from 140° to 200°C, in boreholes in Yellowstone National Park, Wyoming [Bargar and Beeson, 1981; Bargar et al., 1981]. Data from geothermal wells in basalt in Iceland show that phillipsite crystallizes between 65° and 85°C [Kristmannsdottir and Tomasson, 1978].

[11] Group 2 zeolites consist of single 6-ring (Al, Si)<sub>6</sub>O<sub>12</sub> secondary building units in various configurations [*Breck*, 1974]. The only group 2 zeolite included in this study was erionite. It is found in drill cores from rhyolite-derived sediments in geyser basins in Yellowstone National Park, and crystallized at a temperature below  $110^{\circ}$ C [*Honda and Muffler*, 1970]. Data from geothermal wells passing through olivine basalt in Iceland indicate that levyne (which commonly precedes erionite) crystallized in the temperature range from 55° to 70°C [*Kristmannsdottir and Tomasson*, 1978].

[12] Group 4 zeolites consist of double 6-ring (Al, Si)<sub>12</sub>O<sub>24</sub> secondary building units in various configurations [*Breck*, 1974]. Chabazite was the only group 4 zeolite included in this study. Data from geothermal wells in basalt in Iceland show that chabazite crystallizes at depths from 50 to 400 m and at temperatures from 55° to 75°C [*Krist-mannsdottir and Tomasson*, 1978].

[13] Group 5 zeolites consist of complex 4-1,  $T_5O_{10}$  secondary building units in various configurations [*Breck*,

1974]. The group 5 zeolites in this study included natrolite, mesolite, scolecite, and thomsonite. Although natrolite has not been found in active geothermal wells, it should crystallize at temperatures similar to mesolite and scolecite, which are found in geothermal wells in basalt in Iceland, at depths from 450 to 1200 m and at temperatures from 70° to 100°C [*Kristmannsdottir and Tomasson*, 1978]. Liquid inclusions in calcite, found with natrolite on the Keweenaw Peninsula, Michigan, indicate that natrolite crystallized at 133°C [*Stoiber and Davidson*, 1959]. Data from geothermal wells in basalt in Iceland show that mesolite and scolecite both crystallize at depths from 450 to 1200 m and between 70° to 100°C, and thomsonite crystallizes at depths from 100 to 550 m and at temperatures from 65° to 110°C [*Kristmannsdottir and Tomasson*, 1978].

[14] Group 6 zeolites consist of complex 5-1,  $T_8O_{16}$  secondary building units in various configurations [*Breck*, 1974]. The only group 6 zeolite included in this study was mordenite. In Iceland, mordenite is found in basalt, tuffaceous breccia, and dolerite at temperatures from 75° to 280°C and at depths from 25 to 1800 m [*Kristmannsdottir and Tomasson*, 1978]. In the Katayama geothermal area in Japan, mordenite occurs at temperatures from 60° to 140°C [*Seki et al.*, 1969]. In New Zealand, mordenite is found in rhyolite at depths from 118 to 514 m and at temperatures from 60° to 170°C in the Ohaki-Broadlands geothermal area [*Browne and Ellis*, 1970]. It has also been found in pumice and rhyolite tuff taken at depths from 80 to 213 m and at temperatures from 60° to 164°C, in the Wairakei geothermal area [*Steiner*, 1953].

[15] Group 7 zeolites consist of complex 4-4-1,  $T_{10}O_{20}$  secondary building units in various configurations [*Breck*, 1974]. The group 7 zeolites in this study include stilbite, clinoptilolite, heulandite, and stellerite. Data from geothermal wells in basalt in Iceland show that stilbite crystallizes at depths from 45 to 1200 m and at temperatures from 75° to 200°C [*Kristmannsdottir and Tomasson*, 1978]. Clinoptilolite occurs at depths less than 600 m and at temperatures less than 34°C [*Boles*, 1981]. Data from geothermal wells in basalt in Iceland show heulandite crystallizes at depths from 30 to 1200 m and between 65° to 200°C [*Kristmannsdottir and Tomasson*, 1978]. Stellerite is stable to dehydration in a vacuum up to 250°C [*Breck*, 1974].

#### 3. Experimental Procedure

[16] A total of 28 zeolite samples, encompassing 15 different species, were included in this study (Table 1). The samples were crushed by hand in an alumina mortar and pestle and dry sieved to obtain a  $<45 \mu m$  fraction for spectral and compositional analysis. Impurities were removed through a combination of visual inspection with a binocular microscope and with a hand magnet during crushing. Some of the samples were also characterized by powder X-ray diffraction to confirm their identities and to determine whether impurities were present in the samples [*Mertzman*, 2000].

[17] The samples were characterized using X ray fluorescence for major elements [*Mertzman*, 2000] and by instrumental neutron activation analysis (INAA) for selected major elements and minor elements. Ferrous iron was determined using a modification of the procedure outlined by *Reichen and Fahey* [1962]. Ferric iron concentration was taken as the difference between total and ferrous iron. Volatile contents (expressed as loss on ignition in Table 1) were determined by heating splits of the samples to 950°C for 1 hour and measuring the associated weight loss. Sample descriptions, selected major element abundances and volatile content (measured as loss on ignition) are provided in Table 1. Elemental abundances were determined for the volatile-free samples after loss on ignition was measured.

[18] Reflectance spectra for the <45 µm-size fractions were measured at the NASA-supported RELAB spectrometer facility at Brown University. The samples were measured at  $i = 30^{\circ}$  and  $e = 0^{\circ}$  relative to halon for the 0.3- to 2.6- $\mu$ m region at 5 nm spectral resolution and at i = 30° and  $e = 30^{\circ}$  with 4 cm<sup>-1</sup> spectral resolution relative to brushed gold for the 2.5- to 26-µm region. The 0.3- to 2.6-µm spectra were measured under ambient atmosphere. The 2.5to 26-µm spectra were measured after purging the samples (removal of water and CO<sub>2</sub>) for at least 5 hours. Consequently, the 0.3- to 2.5-µm spectra may exhibit spectral features associated with adsorbed water. Details of the RELAB facility and spectral acquisition procedures is given by Pieters [1983] and Reflectance Experiment Laboratory [1996]. The spectra were merged at 2.5  $\mu$ m by multiplying the longer wavelength spectrum by an appropriate correction factor calculated in the region of spectral overlap (2.5-2.6  $\mu$ m). Absorption band depths were calculated using equation (32) of Clark and Roush [1984].

#### 4. Overview of Spectra

[19] The spectral properties of zeolites, both in reflectance and transmittance, have been measured by a number of investigators. The 0.3- to 26-µm reflectance spectra of a number of zeolites are shown in Figure 1. The spectra are broadly similar in nature and are characterized by a number of absorption bands and generally decreasing reflectance toward longer wavelengths.

#### 4.1. Group 1 Zeolites

[20] Compositionally the two wairakite samples are very diverse (Table 1). ZEO105 differs significantly from the ideal formula and that determined for a wairakite from Wairakei, New Zealand [*Deer et al.*, 1963] containing significantly more Si and less Ca. The XRD data indicate that ZEO105 contains both wairakite and quartz. The gismondine, phillipsite and one of the laumontite (ZEO119) samples are compatible with previous compositional determinations and the ideal formulas. The other laumontite sample (ZEO120) contains appreciably more Ca than expected. XRD analysis of this sample indicates that it contains both laumontite and calcite; the presence of calcite would account for the higher-than-expected calcium content.

[21] The spectrum of the <45- $\mu$ m fraction of gismondine is shown in Figure 1. All of the group 1 spectra are characterized by high reflectance in the visible region, >70% at 0.5- $\mu$ m in all cases. The 0.3- to 3.3- $\mu$ m spectra are dominated by absorption features in the 1.4, 1.9, and 2.8- $\mu$ m regions, associated with zeolitic water as discussed below. The absorption features in these regions vary in detail between different minerals. They also exhibit one or two absorption features in the 2.5- to 2.6- $\mu$ m region as

Sample ID <sup>b</sup> MNMH ID <sup>c</sup>	ZEO105 137255	ZEO106 143354	ZEO107 85160	ZEO108 138649	ZEO109 46267-1	ZEO110 83259	ZEO111 93085
Structural group	1	1	1	1	5	3	
SiO <sub>2</sub>	82.04	59.59	46.85	45.36	51.22	46.85	53.63
110 <sub>2</sub>	0.14	0.06	0.00	0.01	0.00	0.00	0.00
$Al_2O_3$ FeO	0.29	23.14	55.00	55.05	51.00	55.01	29.98
FeaOa	0.29	0.91 <sup>e</sup>	0.21 <sup>e</sup>	0 33 <sup>e</sup>	0 18 <sup>e</sup>	$0.12^{e}$	0 11 <sup>e</sup>
MnO	0.029	0.022	0.007	0.008	0.006	0.005	0.005
MgO	0.29	0.22	0.06	0.134	0.084	0.049	0.041
CaO	3.20	13.35	15.28	15.56	11.24	12.32	16.28
Na <sub>2</sub> O	0.151	0.392	0.443	0.45	6.82	6.83	0.24
$\tilde{K_2O}$	2.484	0.015	3.506	2.39	0.009	0.020	0.016
$P_2O_5$	0.030	0.021	0.011	0.019	0.007	0.010	0.007
Sr	80	109	3124	12174	555	3401	90
Zr	98	38	54	<2	30	7	40
V	47	40	35	38	33	32	33
Cr	41	36	39	34	31	33	38
Total <sup>f</sup>	99.52	99.75	99.81	99.36	100.65	99.63	100.34
LOI <sup>S</sup>	2.39	9.01	20.32	21.33	15.15	12.82	14.12
AI.(AI + SI) SCEM <sup>h</sup>	12.5	0.80	44.5	43.7	40.7	42.1	50.0 0.77
	0.90	0.80	0.75	0.74	0.82	0.79	0.77
Sample ID <sup>1</sup>	ZEO112	ZEO113	ZEO114	ZEO115	ZEO116	ZEO117	ZEO118
MNMH ID <sup>c</sup> Structural group <sup>d</sup>	R4303-1 7	113929 5	94550 5	106928 5	162474 5	112868 4	R4175 4
SiO <sub>2</sub>	68.46	51.61	52.12	52.71	50.08	65.30	64.38
TiO <sub>2</sub>	0.00	0.00	0.00	0.00	0.01	0.37	0.00
$Al_2O_3$	19.43	29.35	29.67	29.04	30.25	18.17	22.30
FeO	-	-	-	-	-	0.28	-
Fe <sub>2</sub> O <sub>3</sub>	0.11 <sup>e</sup>	0.10 <sup>e</sup>	0.25 <sup>e</sup>	0.20 <sup>e</sup>	0.11 <sup>e</sup>	1.97	0.14 <sup>e</sup>
MnO	0.004	0.005	0.005	0.010	0.005	0.038	0.013
MgO	0.087	0.134	0.058	0.110	0.051	3.051	0.094
CaO	10.39	10.24	11.05	0.97	1.56	3.40	10.18
Na <sub>2</sub> O PaOa	1.23	8.14	6.68 0.007	17.34	0.040	1.879	1.460
1205	0.000	0.000	0.007	0.007	0.009	0.090	0.007
Sr	77	95	72	115	10201	681	6404
Zr	31	44	49	36	<2	173	<2
V Cr	30	37	33 34	34 122	30	69	32
CI CI	50	35	54	122	55	40	55
Total <sup>r</sup>	99.77	99.73	99.89	100.47	100.16	99.49	100.11
LOI	18.94	13.38	13.00	10.27	10.29	18.39	27.19
Al:(Al + Si)	24.4	39.3	39.3	38.5	40.7	24.0	28.2
SCFM	0.87	0.83	0.82	0.98	0.97	0.91	0.86
Sample ID <sup>j</sup>	ZEO119	ZEO120	ZEO121	ZEO122	ZEO123	ZEO124	ZEO125
MNMH ID <sup>c</sup>	85382	97504	132753	104654	122858	121217	122860
Structural group <sup>a</sup>	1	1	7	7	1	1	2
SiO <sub>2</sub>	65.93	38.09	68.04	67.41	65.35	66.24	70.46
TiO <sub>2</sub>	0.00	0.00	0.00	0.00	0.18	0.66	0.52
$Al_2O_3$	21.30	15.72	20.12	20.76	18.54	18.77	14.81
FeO	- 0.10 <sup>e</sup>	- 0.16 <sup>e</sup>	- 0.12°	_ 	0.22	0.24	- 2.05°
$Fe_2O_3$	0.18	0.10	0.13	0.09	1.92	1.85	3.05
MaO	0.008	0.232	0.008	0.004	0.009	0.018	0.034
CaO	10.61	43.47	7 20	0.020	0.209	1./// 4 70	0.009
Na <sub>2</sub> O	1 10	0.32	3.76	2.00	0.51	1.30	5 20
K <sub>2</sub> O	0.290	0.929	1.062	0.035	4.235	3.471	4.099
$P_2O_5$	0.008	0.031	0.008	0.011	0.071	0.064	0.057
Sr	218	236	723	101	65	2230	287
Zr	23	16	25	23	200	703	577
V	31	36	37	35	60	79	145
Cr	47	38	35	35	39	67	39
Total <sup>f</sup>	99.69	99.43	100.50	100.16	100.35	99.57	100.43
LOI <sup>g</sup>	17.24	26.38	17.60	18.71	16.68	15.02	10.94
Al:(Al + Si)	26.8	31.9	25.1	25.9	24.4	24.4	19.3

 Table 1. Analytical Data for Zeolite Samples<sup>a</sup>

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Table I	(continued)	١.
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Sample ID <sup>j</sup>	ZEO119	ZEO120	ZEO121	ZEO122	ZEO123	ZEO124	ZEO125
MNMH ID <sup>c</sup>	85382	97504	132753	104654	122858	121217	122860
Structural group <sup>d</sup>	1	1	7	7	1	1	2
$SCFM^h$	0.86	0.46	0.90	0.87	0.99	0.91	0.97
Sample ID <sup>k</sup>	ZEO126	ZEO127	ZEO128	ZEO129	ZEO130	ZEO131	ZEO132
MNMH ID <sup>c</sup>	122847	144077	114104-1	122861	122848	R4073-1	138663
Structural group <sup>d</sup>	2	7	7	6	6	7	7
SiO <sub>2</sub>	70.25	73.80	77.37	71.96	64.66	66.46	69.42
TiO <sub>2</sub>	0.32	0.30	0.06	0.68	0.54	0.00	0.00
$Al_2O_3$	15.97	13.22	13.53	14.42	10.00	19.78	19.18
FeO	_	0.24	_	_	_	_	_
Fe <sub>2</sub> O <sub>3</sub>	2.37 <sup>e</sup>	3.01	0.48 <sup>e</sup>	1.93 <sup>e</sup>	4.14 <sup>e</sup>	0.12 <sup>e</sup>	0.15 <sup>e</sup>
MnO	0.053	0.047	0.006	0.013	0.091	0.005	0.006
MgO	1.447	0.337	0.811	0.377	2.555	0.024	0.034
CaO	2.29	1.72	3.31	1.52	10.34	6.66	8.32
Na <sub>2</sub> O	2.18	1.78	1.17	4.10	2.21	2.35	1.78
K <sub>2</sub> O	5.208	4.964	3.325	4.376	4.915	0.942	0.904
$P_2O_5$	0.060	0.040	0.020	0.238	0.233	0.009	0.007
Sr	158	313	154	152	587	29990	4263
Zr	464	565	138	253	174	<2	3
V	116	40	42	331	195	35	32
Cr	40	32	34	39	76	39	34
Total <sup>f</sup>	100.25	99.58	100.13	99.73	99.82	99.91	100.32
LOI <sup>g</sup>	16.14	11.57	13.20	9.41	9.61	15.65	15.70
Al:(Al + Si)	20.5	16.9	16.6	18.5	14.9	25.3	23.9
SCFM <sup>h</sup>	0.95	0.97	0.95	0.97	0.83	0.91	0.89

<sup>a</sup> Values are in wt % for oxides, ppm for elements.

<sup>b</sup> Sample type and location: ZEO105: wairakite, drill hole 66, Wairakei, New Zealand; ZEO106, wairakite, Bandai-atami, Kohriyama, Fukushima, Japan; ZEO107: gismondine, Capo di Bova, Rome, Italy; ZEO108: gismondine, Castel de Decimo, Roma, Lazio, Italy; ZEO109: thomsonite, Elipse Beach, north shore of Lake Superior, Minnesota, USA; ZEO110: thomsonite, North Table Mountain, Golden, Jefferson County, Colorado, USA; ZEO111: scolecite, Hyerabad, India.

<sup>c</sup>Sample identifier used by the National Museum of Natural History of the Smithsonian Institution.

<sup>d</sup>From Breck [1974].

<sup>e</sup>All Fe assumed as Fe<sub>2</sub>O<sub>3</sub>.

<sup>f</sup>Totals are expressed on a volatile-free basis (i.e., elemental abundances determined on sample after measurement of loss on ignition and Sr, Zr, V, and Cr converted to oxides).

<sup>g</sup>Loss on ignition = weight loss after heating sample to 950°C for 1 hour.

<sup>h</sup>SCFM =  $SiO_2/(SiO_2 + CaO + FeO + MgO)$  (for oxide abundances in wt %). Where no FeO abundance is available, all Fe is assumed as Fe<sub>2</sub>O<sub>3</sub>.

<sup>i</sup>Sample type and location: ZEO112: stellerite, Poonah, Maharashtra, India (originally identified as scolecite, a group 5 zeolite); ZEO113: mesolite, Advocate Harbour, Nova Scotia, Canada; ZEO114: mesolite, Cape d'Or, Cumberland County, Nova Scotia, Canada; ZEO115: natrolite, Sao Pedro, Sao Paulo, Brazil; ZEO116: natrolite, Techlovice, Czechoslovakia; ZEO117: chabazite, Christmas, Gila County, Arizona, USA; ZEO118: chabazite, Cape Blomedin, Nova Scotia, Canada.

<sup>j</sup>Sample type and location: ZEO119: laumontite, Great Notch, Essex County, New Jersey, USA; ZEO120: laumontite, Baltimore, Maryland, USA; ZEO121: stilbite, Margaretsville, Nova Scotia, Canada; ZEO122: stilbite, Nova Scotia, Canada; ZEO123: phillipsite, Pine Valley, Eureka County, Nevada, USA; ZEO124: phillipsite, Garbagna, Allesandria, Italy; ZEO125: erionite, Needle Peak, Pershing County, Nevada, USA.

<sup>k</sup>Sample type and location: ZEO126: erionite, Crooked Creek, Rome, Malheur County, Oregon, USA; ZEO127: clinoptilolite, Sheaville, Malheur County, Oregon, USA; ZEO128: clinoptilolite, Fish Creek Mountains, Nevada, USA; ZEO129: mordenite, Trinity Basin, Pershing County, Nevada, USA; ZEO130: mordenite, Crooked Creek, Rome, Malheur County, Oregon, USA; ZEO131: heulandite, Paterson, New Jersey, USA: ZEO132: heulandite, Teigarhorn, Iceland.

either distinct absorption bands or as shoulders superimposed on the short wavelength wing of the O-H stretching fundamental in the 2.8- $\mu$ m region. Differences are also seen in the number and wavelength positions of resolvable Si-O and Al-O stretching bands in the 10- $\mu$ m region and the wavelength position of the Christiansen band in the 8- $\mu$ m region.

#### 4.2. Group 2 Zeolites

[22] Erionite is the only member of this group included in the present study. Compositionally the two samples are consistent with the ideal formula and previous compositional determinations [*Deer et al.*, 1963]. The two erionite spectra are broadly similar to each other as well as to the group 1 zeolites (Figure 1). The longer wavelength bands near 2.8, 6.1, and between 7.7 and 10.1  $\mu$ m are similar between the two samples, with wavelength positions varying by less than 10 nm. Both erionite samples contain a few percent iron whose oxidation state was not determined. Both spectra exhibit weak absorption bands near 0.66 and 0.92  $\mu$ m, suggesting that the iron is present in both ferric and ferrous oxidation states. The spectra also exhibit prominent inflections near 0.37 and 0.49  $\mu$ m, as well as the steep reflectance dropoff shortward of 0.6  $\mu$ m due to metal-oxygen charge transfers [*Burns*, 1993a, 1993b].

#### 4.3. Group 4 Zeolites

[23] This group is represented by two samples of chabazite in the present study. Compositionally the two



**Figure 1.** Reflectance spectra of <45  $\mu$ m grain size fractions of representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1); erionite (ZEO125; group 2); chabazite (ZEO118; group 4); natrolite (ZEO116; group 5); mordenite (ZEO129; group 6); stilbite (ZEO121; group 7): (a) 0.3- to 7- $\mu$ m spectra with each spectrum vertically offset by +0.3 from the spectrum below it for clarity (no offset for the gismondine). (b) 6- to 26- $\mu$ m reflectance spectra of the same samples with each spectrum vertically offset by +0.2 from the spectrum below it for clarity (no offset for the gismondine).

samples are consistent with the ideal formula and those of other chabazites [*Deer et al.*, 1963]. The two chabazite reflectance spectra are similar to each other and broadly similar to the other zeolite spectra (Figure 1). Individual absorption bands in the 2.8- $\mu$ m region are generally unresolved, appearing as a broad region of low reflectance. Absorption bands in the 6.1-, 7.8-, and 10- to 12- $\mu$ m regions show small differences in wavelength positions between the two spectra. The spectra differ from each other shortward of 1.3  $\mu$ m probably due to differences in iron content (Table 1). The higher iron sample (ZEO117) has lower overall reflectance, some evidence for weak ferric and ferrous iron absorptions in the 0.7 and 0.9- $\mu$ m regions, and a steeper dropoff in reflectance shortward of 0.6- $\mu$ m.

#### 4.4. Group 5 Zeolites

[24] This group is represented by two samples each of thomsonite, mesolite, and natrolite, and one sample of scolecite (ZEO111). The two thomsonites, the scolecite (ZEO111), one of the mesolites (ZEO113), and the two natrolite samples are consistent with their ideal compositions and previous compositional determinations [Deer et al., 1963]. One sample which was originally labeled as scolecite (ZEO112) was analyzed using XRD and was found to consist of stellerite with a small amount of stilbite. This interpretation is consistent with the compositional data, and hence this samples was renamed as stellerite, a group 7 zeolite. One of the mesolites (ZEO114) contains more Na than expected (Table 1). The XRD pattern for this sample is more similar to scolecite than mesolite. This is not unexpected as minor substitution of Na for Ca is common in mesolite [Deer et al., 1963]. It should also be noted that zeolite XRD patterns can vary with changes in volatile content [Breck, 1974]. We have chosen to retain the original identity of this sample as mesolite.

[25] Group 5 zeolites, and in particular mesolite, scolecite and natrolite, termed the fibrous zeolites, have been noted by a number of investigators, as possessing transmission spectra which are markedly different from other zeolites [*Milkey*, 1960; *Oinuma and Hayashi*, 1967; *Gottardi and Galli*, 1985]. Group 5 spectra show more complex absorption features than the other groups (Figure 1). The number of resolvable absorption bands in zeolite spectra increases with decreasing openness of the zeolite structure [*Breck*, 1974]. Consequently, the group 5 spectra have the most complex spectra because they have the most compact structures and smallest void spaces.

[26] Natrolite has the most complex spectrum in terms of number of absorption bands (Figure 1) and is similar to natrolite spectra measured by other investigators [e.g., *Grove et al.*, 1992]. This spectral complexity is probably attributable to the higher degree of structural order present in this mineral relative to other group 5 zeolites [*Breck*, 1974]. Water molecules tend to cluster around the Na ions, and the channels in which these clusters reside are small [*Breck*, 1974]. Both effects would tend to lead to more resolved absorption bands as the water molecules are restricted to well-defined local electronic environments. The absorption bands in the 2.8- to 3.2- $\mu$ m region are better resolved than in other zeolite spectra.

#### 4.5. Group 6 Zeolites

[27] This group is represented by two samples of mordenite in the present study. Compositionally the two samples are consistent with the ideal formula and previous analyses [*Deer et al.*, 1963]. One of the mordenite (ZEO129) spectra is shown in Figure 1. The reflectance spectra of the two mordenite samples are similar to each other beyond 1.3  $\mu$ m, but differ shortward of 1.3  $\mu$ m due to differences in iron content and its oxidation state (Table 1). The higher iron sample (ZEO130) has lower overall reflectance and exhibits the iron-related absorption features in the 0.7- and 0.9- $\mu$ m regions and at 0.37 and 0.49  $\mu$ m. At longer wavelengths they are broadly similar to the other zeolite spectra.

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### 4.6. Group 7 Zeolites

[28] This group is represented by two samples each of stilbite, clinoptilolite, and heulandite, and one sample of stellerite. All of these samples are consistent with their ideal formulas and previous compositional determinations [*Deer et al.*, 1963]. The reflectance spectrum of a stilbite (ZEO121) is shown in Figure 1.

[29] The group 7 spectra show less diversity than the group 5 spectra. They do not show well-resolved absorption bands in the 2.8-µm region. The H-O-H bending fundamental also shows little variation within this group, varying in wavelength position from 6.11 to 6.15 µm. All of the group 7 spectra exhibit three resolvable absorption features in the 7.5- to 11-µm region. There is more spectral variability in this region, with the wavelength position of the Christiansen band ranging between 7.67 and 7.87 µm. All of the spectra exhibit an absorption feature near 2.55  $\mu$ m and two partially overlapping absorption bands each in the 1.4 and 1.9-µm regions. A number of the spectra also exhibit absorption bands near 0.97 and 1.16 µm. The spectrum of the most iron-rich sample (ZEO127) has the lowest overall reflectance shortward of 1.3 µm, weak but resolvable ferric and ferrous iron absorption bands in the 0.7 and 0.9-µm regions, and the steepest reflectance dropoff shortward of 0.6 µm.

### 5. Spectral-Compositional-Structural Relationships

[30] Overall, the zeolites included in this study exhibit the same broad types of absorption features; below  $\sim$ 7 µm, the spectra are dominated by absorption features in the 1.4-, 1.9-, 2.5-, 2.8-, and 6.1-µm regions. Longward of this wavelength the spectra are characterized by low overall reflectance and common sets of absorption features in various wavelength regions. In the ensuing discussion, we have grouped spectral features on the basis of the major compositional parameter associated with each.

[31] The absorption features in the zeolite reflectance spectra, particularly beyond  $\sim 8 \,\mu\text{m}$ , are much broader than comparable absorption bands in transmission spectra. This wavelength region is also characterized by low overall reflectance. Thus, the various absorption bands beyond 8  $\mu\text{m}$  are intense, and their broadness results in low overall reflectance and a loss of spectral detail. Similar results were found in a recent comparison of diaspore reflectance and transmission spectra [*Cloutis and Bell*, 2000]. These differences are attributable to the different scattering mechanisms inherent in transmission spectral studies have provided important insights into the causes of various absorption features.

#### 5.1. Iron-Related Spectral Features

[32] A number of the zeolite samples contain a few wt % FeO and/or Fe<sub>2</sub>O<sub>3</sub> (Table 1). This results in absorption features near 0.7 and/or 0.9  $\mu$ m (Figure 2) which are

**Figure 2.** 0.3- to 1.3- $\mu$ m reflectance spectra of various iron-bearing zeolites showing evidence for iron-related absorption features near 0.37, 0.46, 0.7, and 0.9  $\mu$ m. Each spectrum has been vertically offset by +0.2 from the one below it for clarity. From bottom to top: phillipsite (group 1; ZEO123); erionite (group 2; ZEO126); mordenite (group 6; ZEO130); clinoptilolite (group 7; ZEO127).

attributable to an Fe<sup>3+</sup>-Fe<sup>2+</sup> intervalence charge transfer and an Fe<sup>2+</sup> crystal field transition, respectively [e.g., *Burns*, 1993a, 1993b]. The high overall reflectance of the spectra allows these bands to be resolved in spite of the relatively low iron contents. The spectra with Fe-associated absorption features also exhibit the steepest reflectance decreases shortward of 0.6  $\mu$ m, which is attributable to Fe-O charge transfers [*Burns*, 1993a, 1993b].

[33] The spectra of the samples with the highest Fe contents also exhibit weak absorption features near 0.37 and 0.49  $\mu$ m which are attributed to Fe<sup>2+</sup> crystal field transitions [*Burns*, 1993a, 1993b]. The presence of iron does not appear to have a systematic effect on the wavelength positions of the various OH-associated absorption features in the 1.4 and 1.9- $\mu$ m regions, although it is possible that small wavelength shifts in band positions could be offset by structural differences or other cation substitutions. There is an additional absorption feature associated with iron and water at 2.30  $\mu$ m which is discussed below. Table 2 summarizes the assignments and compositional or structural information associated with the various absorption bands seen in the zeolite spectra.

#### 5.2. Water-Related Spectral Features

[34] A number of investigators have shown that water contained within the voids of the zeolite structure (zeolitic water) is the main contributor to absorption bands associated with  $H_2O$  stretching and bending vibrations; OH and adsorbed water are relatively insignificant spectrally



Wavelength, µm	Assignment	Notes
	Groups 1, 2, 4, 6, and 7 Zeolites and T	Thomsonite
0.46	Fe <sup>2+</sup> crystal field transition	Present in Fe <sup>2+</sup> -bearing zeolites
$\sim 0.7$	$Fe^{3+}-Fe^{2+}$ intervalence charge transfer	Present in Fe <sup>3+</sup> /Fe <sup>2+</sup> -bearing zeolites
$\sim 0.9$	Fe <sup>2+</sup> crystal field transition	Present in Fe <sup>2+</sup> -bearing zeolites
0.97	Second overtone of H <sub>2</sub> O stretch	Present across all classes
1.155 - 1.160	Symmetric + asymmetric $H_2O$ stretch + $H_2O$ bend (bound water)	Na- and K-rich zeolites
1.160 - 1.180	Symmetric + asymmetric $H_2O$ stretch + $H_2O$ bend (bound water)	Ca-rich zeolites
1 19	Symmetric + asymmetric $H_2O$ stretch + $H_2O$ bend (obtaind water)	Present across all classes
1.417 - 1.422	$H_2O$ stretch + first overtone of $H_2O$ bend (bound water)	Na- and K-rich zeolites
1.424 - 1.430	$H_2O$ stretch + first overtone of $H_2O$ bend (bound water)	Ca-rich zeolites
~1.47	$H_2O$ stretch + first overtone of $H_2O$ bend (obtaind water)	Present across all classes
1 911	$H_2O$ stretch + hend (bound water)	K-rich zeolites
1 913	$H_2O$ stretch + bend (bound water)	Na-rich zeolites
1 922	$H_2O$ stretch + bend (bound water)	Ca-rich zeolites
$\sim 1.922$	$H_2O$ stretch + bend (adsorbed water)	Present across all classes
2 30	$H_2O$ stretch + bend (AlFe-OH)	Present across all classes
2.50 2 460 - 2 465	$H_2O$ stretch + bend (AlCa-OH)	Present across all classes
$\sim 2.400$ 2.405 $\sim 2.80$	H <sub>2</sub> O stretch (bound water)	Present across all classes
~3.07	First overtone of H <sub>2</sub> O bend (bound water)	Present across all classes
6.05 - 6.15	$H_2O$ bend (bound water)	Present across all classes
7.50 - 8.53	Christiansen frequency	Correlated with $A1:(A1 + Si)$ ratio
83-94	Asymmetric Si-O/Al-O stretch	Correlated with $A1:(A1 + Si)$ ratio
94 - 107	Asymmetric Si-O/Al-O stretch	Correlated with $A1:(A1 + Si)$ ratio
112 - 121	Transparency peak	Correlated with $A1:(A1 + Si)$ ratio
$\sim 12.5 - 15$	Symmetric Si-O/Al-O stretch	Some structural group discrimination
$\sim 20 - 26$	O-Si-O/O-Al-O bends	Discrimination of group 5 zeolites
	Group 5 Fibrous Zeolites (Water-Related Abs	sorption Bands)
1.427 - 1.430	H <sub>2</sub> O stretch + first overtone of H <sub>2</sub> O bend (Ca-bound water)	Position increases and depth decreases with increasing Na
1.436	$H_2O$ stretch + first overtone of $H_2O$ bend (Ca-bound water)	Only present in scolecite
1.445 - 1.458	$H_2O$ stretch + first overtone of $H_2O$ bend (Na-bound water)	Position and depth increase with increasing Na
$\sim 1.47$	$H_2O$ stretch + first overtone of H2O bend (absorbed water)	Partially obscured by other bands
1.485 - 1.489	$H_2O$ stretch + first overtone of $H_2O$ bend (Ca-bound water)	Position increases and depth decreases with increasing Na
1.940 - 1.945	$H_2O$ stretch + bend (Ca-bound water)	Depth decreases with increasing Na
$\sim 1.970$	$H_2O$ stretch + bend (adsorbed water)	Present across all classes
2.005	$H_2O$ stretch + bend (Na-bound water)	Depth increases with increasing Na
2.080	$H_2O$ stretch + bend (Na-bound water)	Depth increases with increasing Na
2.175	$H_2O$ stretch + bend (AlCa-OH)	Depth increases with increasing Ca
2.200	$H_2O$ stretch + bend (AlNa-OH)	Depth increases with increasing Na
2.405	$H_2O$ stretch + bend (AlCa-OH)	Depth increases with increasing Ca
2.430	$H_2O$ stretch + bend (AlNa-OH)	Depth increases with increasing Na
2.465	$H_2O$ stretch + bend (AlCa-OH)	Depth increases with increasing Ca
$\sim 2.54$	$H_2O$ stretch + bend (AlCa-OH)	Depth increases with increasing Ca
2.590	$H_2O$ stretch + bend (AlNa-OH)	Depth increases with increasing Na
2.790	$H_2O$ stretch (Ca-bound water)	Depth increases with increasing Ca
2.825	$H_2O$ stretch (Na-bound water)	Depth increases with increasing Na
2.925	$H_2O$ stretch (Ca-bound water)	Depth increases with increasing Ca
3.00	$H_2O$ stretch (Na-bound water)	Depth increases with increasing Na
3.09	$H_2O$ stretch (Ca-bound water)	Depth increases with increasing Ca
3.12	$H_2O$ stretch (Na-bound water)	Depth increases with increasing Na
6.01	H <sub>2</sub> O bend (Ca-bound water)	Depth increases with increasing Ca
6.03	H <sub>2</sub> O bend (Na-bound water)	Depth increases with increasing Na
6.06	$H_2O$ bend (Ca-bound water)	Depth increases with increasing Ca
6.12	$H_2O$ bend (Na-bound water)	Depth increases with increasing Na
6.28	H2O bend (Ca-bound water)	Depth increases with increasing Ca

 Table 2.
 Absorption Band Assignments for Zeolites

[*Bertsch and Habgood*, 1963; *Angell and Schaffer*, 1965]. Hence the ensuing discussion relates exclusively to zeolitic water unless otherwise indicated.

#### 5.2.1. 0.97-µm Region

[35] A number of zeolite reflectance spectra exhibit an absorption feature in the 0.97- $\mu$ m region (Figure 3) [*Hunt and Salisbury*, 1970; *Grove et al.*, 1992]. This feature is attributed to the second overtone of the O-H stretch [*Hunt and Salisbury*, 1970; *Clark et al.*, 1990] and is most evident in the reflectance spectra with high overall reflectance in this region. Its depth is on the order of 2% at most, as expected for a second order overtone. It appears in too few

of the zeolite spectra, and is not sufficiently well-resolved in the current data to permit a more detailed analysis. This band may be due to either bound or adsorbed water, as the spectra in this region were not measured under dry conditions.

#### 5.2.2. 1.16-µm Region

[36] A number of the spectra also exhibit an absorption feature near 1.165  $\mu$ m (Figure 3). This band appears in most of the spectra as a weak (<2% D<sub>b</sub>), but distinct, narrow feature. It can be attributed to a combination of a symmetric O-H stretch, an asymmetric O-H stretch and the H-O-H bending fundamental [*Hunt and Salisbury*, 1970; *Clark et* 



**Figure 3.** 0.8- to 1.3- $\mu$ m reflectance spectra of various zeolites showing evidence for H<sub>2</sub>O-related absorption features in the 0.97 and 1.16- $\mu$ m regions. From bottom to top: laumontite (ZEO119; group 1); chabazite (ZEO118; group 4); scolecite (ZEO112; group 5); stilbite (ZEO121; group 7).

al., 1990]. This band is only associated with the presence of water and not OH. It is most commonly found in the spectra of the most volatile-rich zeolites, as expected (groups 1, 2, and 7; Table 1). This feature consists of a strong band in the 1.155- to 1.180-µm region and a weaker shoulder near 1.19  $\mu$ m. We assign the former to bound water and the latter to adsorbed water, analogous to the results found by Bishop et al. [1994] for montmorillonite, where bound water absorption bands occur at shorter wavelengths than adsorbed water absorption bands. The wavelength position of the 1.16-µm band shows some correlation with the major cation present. Its wavelength position generally increases as the major cation weight increases. In Na- and K-rich zeolites, the band occurs between 1.155 and 1.160 µm, while in Ca-rich zeolites it occurs between 1.160 and 1.180 µm. It occurs at generally longer wavelengths in those zeolites with larger voids, consistent with less hydrogen bonding to cations expected in such zeolites (Table 2).

### 5.2.3. 1.4-µm Region

[37] All of the zeolite spectra exhibit absorption features in the 1.4- $\mu$ m region. For the majority, this feature consists of a stronger absorption band near 1.42- $\mu$ m and a weaker band (often appearing as a shoulder) near 1.47  $\mu$ m (Figure 4). The 1.42- $\mu$ m band is attributed to bound H<sub>2</sub>O combination bands, while the 1.47- $\mu$ m band is assigned to adsorbed H<sub>2</sub>O combination bands [*Bishop et al.*, 1994] because its wavelength position is largely invariant (1.465–1.470  $\mu$ m). These features involve an H<sub>2</sub>O stretch plus the first overtone of the H<sub>2</sub>O bend. If group 5 zeolites are ignored, we observe a general increase in the wavelength position of the main



**Figure 4.** 1.2- to 1.6- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites showing details of absorption features in the 1.4- $\mu$ m region. Some spectra have been vertically offset by the amount indicated in brackets for clarity. (a) Representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1; no offset); erionite (ZEO125; group 2; +0.05); chabazite (ZEO118; group 4; +0.2); natrolite (ZEO116; group 5; +0.3); mordenite (ZEO129; group 6; +0.3); stilbite (ZEO121; group 7; +0.4). (b) The same for group 5 zeolites; from bottom to top: thomsonite (ZEO109; no offset); natrolite (ZEO116; +0.02); scolecite (ZEO111; +0.06); mesolite (ZEO113; no offset).



**Figure 5.** 1.7- to 2.2- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites showing details of absorption features in the 1.9- $\mu$ m region. (a) Representative zeolites from each structural group. Each spectrum has been vertically offset by +0.3 from the spectrum below it for clarity. From bottom to top: gismondine (ZEO107; group 1; no vertical offset); erionite (ZEO125; group 2); chabazite (ZEO118; group 4); natrolite (ZEO116; group 5); mordenite (ZEO129; group 6); stilbite (ZEO121; group 7). (b) The same for group 5 zeolites (vertical offsets are indicated in brackets); from bottom to top: thomsonite (ZEO109; no offset); mesolite (ZEO113; no offset); scolecite (ZEO111; +0.3); natrolite (ZEO116; +0.4).

absorption band from 1.420  $\mu$ m in K-bearing zeolites, to 1.422  $\mu$ m in Na-bearing zeolites to 1.427  $\mu$ m in Ca-bearing zeolites (Table 2). Thus, the wavelength position of this absorption band can be used to distinguish the major cation present in zeolites. These differences are attributed to the variations in the degree of bonding with the various cations [*Bishop et al.*, 1994]. The different structural groups do not exhibit any systematic trends in terms of positions of these absorption bands.

[38] The group 5 zeolites scolecite, thomsonite, mesolite, and natrolite exhibit more complex absorptions in this wavelength region (Figure 4). Structurally, group 5 zeolites are distinguished from other zeolites by their small channels and clustering of water molecules around the cations. This, coupled with the presence of both Ca and Na in many members of the group (Table 1) should give rise to at least two absorption bands in the 1.4-µm region in addition to the adsorbed water band. This can be seen by comparing the spectra of the scolecite, mesolite, and natrolite (Figure 4). All three are structurally similar with different Na:Ca ratios (Table 1). As Na content increases, the absorption bands in the 1.427- to 1.430- and 1.485- to 1.489-µm regions move to longer wavelengths and decrease in intensity, the bands near 1.436 µm become obscured (present only in the scolecite spectrum), and the band in the 1.445- to 1.458µm region moves to longer wavelengths and becomes deeper. These results suggest that the 1.436- and 1.445- to 1.458-µm bands are associated with Na-bonded water, while the 1.427- to 1.430- and 1.485- to 1.489- $\mu$ m bands are associated with Ca-bonded water in these zeolites. It should be noted that the group 5 fibrous zeolites can undergo irreversible dehydration, accompanied by structural rearrangements, and hence the absorption bands in this region are only characteristic of the unaltered minerals.

#### 5.2.4. 1.9-µm Region

[39] Absorption bands in the 1.9-µm region are attributable to a combination of the H<sub>2</sub>O stretching and bending modes [Clark et al., 1990; Bishop et al., 1994]. As was the case for the 1.4-µm region, the majority of the zeolite spectra, with the exception of group 5, exhibit a strong absorption band near 1.92 µm and a weaker band near 1.97 µm (Figure 5). The strong band is attributed to bound H<sub>2</sub>O combination bands, while the 1.97-µm band is assigned to adsorbed H<sub>2</sub>O combination bands [Bishop et al., 1994]. The wavelength position of the band near  $1.92 \text{-}\mu\text{m}$ ranges from 1.911 to 1.930  $\mu$ m. The same trends as was observed for the 1.4-µm band is also observed here; i.e., the position of this band generally increases from K-bearing zeolites (1.911 µm average) to Na-bearing zeolites (1.913  $\mu$ m average) to Ca-bearing zeolites (1.922  $\mu$ m average) (Table 1). The position of the band near 1.97 µm is largely invariant as would be expected for adsorbed water, ranging between 1.970 and 1.975 µm. It is present in spectra from all the structural groups.

[40] The group 5 fibrous zeolite spectra are more complex in this wavelength region, while thomsonite is similar



**Figure 6.** 2.1- to 2.75- $\mu$ m zeolite reflectance spectra showing details of various cation-H<sub>2</sub>O absorption features. Some spectra have been vertically offset by the amount indicated in brackets for clarity. Ca-related absorption band near 2.465  $\mu$ m is present in gismondine (ZEO108; lower spectrum; no offset) and wairakite (ZEO116; second spectrum from bottom; no offset). An Fe-related absorption band near 2.3  $\mu$ m is present in clinoptilolite (ZEO127; third spectrum from bottom; +0.2) and erionite (ZEO125; top spectrum; +0.3).

to the other zeolites described above. Scolecite and mesolite are somewhat similar to each other, with well-resolved absorption bands in the 1.940- to 1.945- and 1.975- to 1.977-µm regions, but they also exhibit additional less wellresolved bands at longer wavelengths. Natrolite exhibits the most unique spectrum in this region, with well-resolved absorption bands at 1.942, 2.005, and 2.080 µm (Figure 5) [*Grove et al.*, 1992]. The depth of the band at 1.940–1.945 µm decreases with increasing Na content and there is no clear trend involving band position and Na:Ca ratio. The depths of the bands at 2.005 and 2.080 µm both increase with increasing Na content and are attributed to the Nawater clusters in natrolite [*Breck*, 1974].

#### 5.2.5. 2.1- to 2.5-µm Region

[41] Approximately half of the zeolites in this study exhibit absorption bands in the 2.1- to 2.5-μm region (Figure 6). Absorption band seen in this wavelength region for H<sub>2</sub>O/OH-bearing silicates have traditionally been assigned to combinations of the OH stretching and bending vibrations for metal-OH groups. [*Stubičan and Roy*, 1961; *Freund*, 1974; *Moenke*, 1974; *Clark et al.*, 1990]. OH stretching vibrations occur in the 2.8-μm region, while H-O-H librations occur in the 10- to 30-μm region [*Bayly et al.*, 1963; *Ryskin*, 1974], thus absorption bands are expected in the 2.1- to 2.5-μm region due to combinations of these modes, with the wavelength position of the combination bands being a function of the particular cation to which the H<sub>2</sub>O is hydrogen bonded. In the case of the current zeolites, the main metals associated with these bands are Ca, Na, K, and Fe. While OH is not expected in zeolites [Bertsch and Habgood, 1963], a certain amount of hydrogen bonding can occur between the H<sub>2</sub>O hydrogen atoms and the framework oxygen atoms as well as with other cations [Bertsch and Habgood, 1963; Angell and Schaffer, 1965; Breck, 1974; Ryskin, 1974]. Three main groups of absorptions were found. The iron-bearing zeolites display a weak absorption feature near 2.30 µm, the Cabearing zeolites show an absorption feature near 2.465 µm, and the Na- and K-bearing zeolites display an absorption feature near 2.68  $\mu$ m. We attribute these features to H<sub>2</sub>O stretching plus bending vibrations involving AlFe-OH, AlCa-OH, and Al(Na,K)-OH molecular clusters, respectively. These differences are consistent with the shifts in band positions as a function of a particular cation found by *Bishop* et al. [1994] in montmorillonite spectra.

[42] Once again, the group 5 zeolites exhibit the most complex and numerous absorption features in this wavelength region (Figure 6). On the basis of relative band depths and positions, we assign absorption bands at 2.175, 2.405, and 2.465  $\mu$ m to AlCa-OH absorptions, as all of these features become more intense as Ca content increases. The bands at 2.175 and 2.465  $\mu$ m are also seen, weakly, in some of the nonfibrous Ca-rich zeolite spectra, lending further support to these assignments. Bands at 2.200, 2.430, and 2.590 m are probably associated with NaAl-OH absorptions because their intensity increases with Na content.

#### 5.2.6. 2.6- to 3.2-µm Region

[43] Unlike the shorter wavelength regions, the spectra beyond 2.5  $\mu$ m were measured under dry atmospheric (purged) conditions. Thus, we expect adsorbed water to contribute little if anything to the spectra in this region. The spectral complexity seen at shorter wavelengths, particularly in the 1.4, 1.9, and 2.0- to 2.5- $\mu$ m regions is mirrored in the 3- $\mu$ m region. As an example, the group 5 zeolites, which commonly exhibit multiple absorption bands in the 1.4 and 1.9- $\mu$ m regions, exhibit the greatest number of absorption bands in the 3- $\mu$ m region (Figure 7).

[44] The 2.6- to 3.2-µm region is characterized in all cases by a series of intense absorption bands. These bands are associated with symmetric and asymmetric O-H stretching vibrations ( $\nu_1$  and  $\nu_3$ ) as well as the first overtone of the H-O-H bending fundamental ( $\nu_2$ ) in zeolitic water [*Milkey*, 1960; Saksena, 1961; Bertsch and Habgood, 1963; Angell and Schaffer, 1965; Oinuma and Hayashi, 1967; Moenke, 1974; Gottardi and Galli, 1985; Tsitsishvili et al., 1992]. The number, wavelength position, and intensity of these bands varies as a function of many factors, such as the degree of cation-H<sub>2</sub>O or structural oxygen-H<sub>2</sub>O bonding, nature of the cations, water content, and electronic environment of the sites occupied by the water molecules [Bertsch and Habgood, 1963; Bayly et al., 1963; Angell and Schaffer, 1965; Oinuma and Hayashi, 1967; Breck, 1974; Ryskin, 1974]. Differences in transmission spectra in this wavelength region have been used to distinguish different zeolite structural groups [e.g., Oinuma and Hayashi, 1967].

[45] All of the reflectance spectra exhibit a broad absorption feature near 2.6  $\mu$ m, which appears as a prominent



**Figure 7.** 2.6- to 3.2- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites. Some spectra have been vertically offset by the amount indicated in brackets for clarity. (a) Representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1; no offset); erionite (ZEO125; group 2; +0.1); chabazite (ZEO118; group 4; +0.3); natrolite (ZEO116; group 5; +0.4); mordenite (ZEO129; group 6; +0.6); stilbite (ZEO121; group 7; +0.9). (b) The same for group 5 zeolites. From bottom to top: thomsonite (ZEO109; no offset); scolecite (ZEO111; +0.06); mesolite (ZEO113; +0.1); natrolite (ZEO116; +0.2).

inflection on the short wavelength wing of the fundamental H<sub>2</sub>O stretching band (Figure 6). This band is assigned to an asymmetric stretch in H<sub>2</sub>O ( $\nu_3$ ) plus the H<sub>2</sub>O libration ( $\nu_L$ ) for bound water [*Bayly et al.*, 1963]. An absorption band in this region appears in reflectance spectra of other minerals containing "free" water such as beryl, as well as in H<sub>2</sub>O transmission spectra [*Bayly et al.*, 1963; *Clark et al.*, 1990].

[46] Most of the zeolite spectra are characterized by a broad region of intense absorption, with minima near 2.8 and 3.07  $\mu$ m. These features are assigned to H<sub>2</sub>O stretching vibrations and the first overtone of the H<sub>2</sub>O bending vibration, respectively [Bertsch and Habgood, 1963; Bavlv et al., 1963; Angell and Schaffer, 1965; Oinuma and Hayashi, 1967; Breck, 1974; Ryskin, 1974; Clark et al., 1990; Bishop et al., 1994]. There is generally insufficient resolution of individual absorption bands in this region to clearly resolve symmetric from asymmetric stretches. Previous investigators have found small differences in terms of the wavelength positions of these bands as a function of cation type and zeolite structural group [Angell and Schaffer, 1965] in transmission spectra, with the wavelength position of these bands moving to longer wavelengths as the degree of hydrogen bonding increases [Oinuma and Hayashi, 1967]. The current spectra do show a difference in the position of the band near 2.8 µm; it occurs near 2.825 µm in the group 5 spectra, and generally near 2.80 µm in the others. This is consistent with previous interpretations, where hydrogen bonding is greatest in group 5 zeolites [Tsitsishvili et al., 1992]. The intensity and broadness of these absorption features in most of the zeolite reflectance spectra precludes more detailed analysis in terms of correlating small changes in wavelength positions with either structural or compositional variations. Band depths in this region can reach 95% with as little as 10 wt % volatiles (presumed to be water).

[47] The group 5 spectra, once again, show the greatest spectral complexity in this wavelength region (Figure 7) [Oinuma and Hayashi, 1967]. They are characterized by a number of absorption features which vary from sample to sample. On the basis of these differences and in conjunction with the other zeolite spectra, we assign the absorption feature at 2.790 and 2.925, and 3.09  $\mu$ m to Ca-bound H<sub>2</sub>O stretching vibrations, and the bands at 2.825 and 3.00, and 3.12  $\mu$ m to Na-bound H<sub>2</sub>O stretching vibrations. The absorption band at 3.09 µm, most prominent in the Ca-rich group 5 spectra, is assigned to the first overtone of the  $H_2O$ bending vibration for Ca-rich group 5 zeolites, and the absorption band at 3.00 µm, most prominent in the Na-rich group 5 spectra, is assigned to the first overtone of the  $H_2O$ bending vibration for Na-rich group 5 zeolites. As expected given its intermediate composition, the mesolite shows evidence for all of these features. Again, we attribute the multiplicity of bands in the group 5 spectra to the fact that water is highly confined in the channels in these zeolites and can form clusters with the various cations, leading to more discrete energy levels, and hence more resolvable absorption bands [Breck, 1974].

#### 5.2.7. 6.1-µm Region

[48] All of the zeolite spectra exhibit an absorption feature near 6.1  $\mu$ m. This feature is attributed to H-O-H bending vibrations in H<sub>2</sub>O ( $\nu_2$ ) [*Clark et al.*, 1990; *Bishop et al.*, 1994]. Given that water is the dominant form of



**Figure 8.** 5.8- to 6.8- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of selected zeolites. Some spectra have been vertically offset by the amount indicated in brackets for clarity. From bottom to top: chabazite (ZEO118; group 4; no offset); stilbite (ZEO122; group 7; +0.04); laumontite (ZEO119; group 1; +0.05); scolecite (ZEO111; group 5; +0.1); natrolite (ZEO116; group 5; +0.16).

hydroxyl in zeolites, we see an intense absorption feature in this wavelength region. In the case of the zeolites with only a single absorption band in this region, it occurs at  $6.1 \pm 0.05$  $\mu$ m (Figure 8). Again, the group 5 spectra exhibit more complex behavior in this region with differences in wavelength positions associated with different cations and degree of hydrogen bonding, but with all bands attributable to H<sub>2</sub>O bending vibrations [*Angell and Schaffer*, 1965; *Oinuma and Hayashi*, 1967]. The presence of water molecules in discrete configurations surrounding the various cations in the confined sites and the associated hydrogen bonding leads to complex spectral behavior. The bands at 6.01, 6.06, and 6.28  $\mu$ m are attributed to Ca-bound H<sub>2</sub>O bends, and the bands at 6.03 and 6.12  $\mu$ m are attributed to Na-bound H<sub>2</sub>O bands.

#### 5.2.8. Water Band Summary

[49] As is evident from the water-associated absorption features, the number and position of these bands is a function of a combination of both structure and composition. In the case of zeolites with large void spaces, water-related absorption bands are generally few in number, broad, and relatively constant in wavelength position. In zeolites with small void spaces, hydrogen bonding of water molecules to either cations in the void spaces or oxygens which are part of the aluminosilicate framework is enhanced. The presence of multiple and energetically well-constrained environments for water molecules will lead to multiple and well-resolved water-associated absorption bands.

#### 5.3. Si- and Al-Related Spectral Features

[50] The aluminosilicate framework of zeolites, consisting of different configurations of  $SiO_4$  and  $AlO_4$  tetrahedra, can give rise to absorption features at generally longer

wavelengths than those associated with zeolitic water. As was the case with water, such absorption features can be attributed to stretching and bending vibrations whose energy can be mediated by neighboring cations and structural variations which alter the local electronic environment. **5.3.1. 4-** to **5-\mum Region** 

[51] A number of the zeolite spectra, and in particular those of group 5, exhibit an absorption feature in the 4.4- to 4.9- $\mu$ m region (Figure 9). These features are probably the first overtone of the main Si-O stretching vibration present in the 9- to 10- $\mu$ m region (see below), as changes in the wavelength position and depths of the absorption bands in the 4.4- to 4.9- $\mu$ m region closely follow those in the 9- to 10- $\mu$ m region.

#### 5.3.2. Christiansen Frequency

[52] The most intense absorption band is almost always present between 7.50 and 8.53  $\mu$ m, similar to the reflectance spectra of zeolites measured by other investigators (Figure 9) (e.g., Atmosphere-Surface Turbulent Exchange Facility (ASTER). Web site: http://www.speclib.ipl.nasa.gov. 2000). This band is termed the Christiansen frequency and it occurs where the refractive index of the mineral passes through unity; it is present on the short wavelength wing of an absorption band [Logan et al., 1973; Salisbury, 1993]. The wavelength position of the Christiansen frequency is positively correlated with the Al:(Al + Si) ratio (Figure 10) with one exception (ZEO130, mordenite). The higher than expected value for this sample may be due to the intense fundamental band located at longer wavelengths which skews the Christiansen band minimum toward longer wavelengths. This correlation permits the Al:(Al + Si)



**Figure 9.** 3.4- to 5.0- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of selected zeolites which exhibit resolvable absorption bands in this region. Some spectra have been vertically offset by the amount indicated in brackets for clarity. From bottom to top: stilbite (ZEO122; group 7; no offset); chabazite (ZEO118; group 4; no offset); laumontite (ZEO119; group 1; +0.05); scolecite (ZEO111; group 5; +0.1); natrolite (ZEO116; group 5; +0.2).



**Figure 10.** Wavelength position of the Christiansen band in the 7.5- to 8.5-µm region of the zeolite reflectance spectra versus Al:(Al + Si) ratio.

ratio for zeolites to be rapidly constrained from their reflectance spectra. However, it should be noted that the wavelength position of this band can vary as function of viewing geometry, grain size, particle packing, and atmospheric pressure [Logan et al., 1973; Salisbury and D'Aria, 1989; Salisbury and Walter, 1989], all of which are relatively constant in this study.

[53] The wavelength position of the Christiansen frequency has also been found to be useful for more complex compositional determinations, such as the SCFM index  $(SiO_2/(SiO_2 + CaO + FeO + MgO))$  [Salisbury and Walter, 1989]. We found a weak inverse trend between the SCFM index and the wavelength position of the Christiansen band with a fair degree of scatter, similar to the results of *Salisbury and Walter* [1989]. The amount of scatter was much greater than for the Al:(Al + Si) ratio determination. The caveats described above also apply to this determination.

#### 5.3.3. 8.5- to 11-µm Region

[54] Previous transmission spectral studies of zeolites have shown that these minerals exhibit a number of absorption bands in the 8.5- to 11-µm region, which vary among different zeolite structural groups [e.g., Oinuma and Hayashi, 1967; Breck, 1974]. These bands are attributed to asymmetric O-Si-O and O-Al-O stretching fundamentals (restrahlen bands). However, beyond attributing the most intense of these bands to Si-O stretches, there is generally little agreement on more precise band assignments [Milkey, 1960; Saksena, 1961; Stubičan and Roy, 1961; Oinuma and Hayashi, 1967; Breck, 1974; Moenke, 1974; Gottardi and Galli, 1985; Salisbury and Walter, 1989; Tsitsishvili et al., 1992]. They are the most intense absorption bands observed in zeolite transmission spectra, and are generally quite broad and few in number [Milkey, 1960; Oinuma and Hayashi, 1967; Gottardi and Galli, 1985]. Spectral differences among zeolites in this wavelength region are expected as the number and wavelength position of these bands will be affected by factors such as Si and Al abundances and structure [e.g., Milkey 1960]. Si-Al substitution causes a sufficiently large change in the nature of the stretching vibrations that the wavelength position of the absorption peak in the 9- to 10-µm region in transmission spectra can be correlated with Al:(Al + Si) ratio for many tectosilicates [Milkey, 1960]. This peak position ranges from 9.18 m in



**Figure 11.** 7- to 13- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites showing details of absorption features in this region. Some spectra have been vertically offset by the amount indicated in brackets for clarity. (a) Representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1; no vertical offset); erionite (ZEO125; group 2; +0.1); chabazite (ZEO118; group 4; +0.2); natrolite (ZEO116; group 5; +0.3); mordenite (ZEO129; group 6; +0.5); stilbite (ZEO121; group 7; +0.6). (b) The same for group 5 zeolites; from bottom to top: thomsonite (ZEO109; no offset); scolecite (ZEO111; +0.06); mesolite (ZEO113; +0.15); natrolite (ZEO116; +0.2).



**Figure 12.** Al:(Al + Si) ratio versus wavelength positions of the two major O-Si-O/ O-Al-O asymmetric stretching fundamentals located in the 8.2- to 9.4-µm (squares) and 9.4- to 10.7-µm (circles) regions of the zeolite reflectance spectra.

quartz (Al:(Al + Si) ratio of 0) to 10.0 µm (Al:(Al + Si) ratio of 0.5). This trend is apparently insensitive to cation substitutions [Milkey, 1960].

[55] The 8.5- to 11- $\mu$ m region of the zeolite reflectance spectra generally show two major absorption bands in the 8.3- to 9.4- and 9.4- to 10.7-μm region (Figure 11). The fibrous zeolite spectra generally exhibit at least 5 bands in this same region. Comparable trends to those found by Milkev [1960] were found in our reflectance spectra for both the short (8.3-9.4) and long (9.4-10.7) wavelength bands (Figure 12), indicating that the wavelength positions of these bands in reflectance spectra are useful for constraining Al:(Al + Si) ratios. There is overlap between the different structural groups, suggesting that these band positions are more sensitive to Al:(Al + Si) ratios than structure. However, the positions of these bands is useful for discriminating at least some of the structural groups (Table 3). Within each of the groups there is also a positive correlation between the position of both of these bands and Al:(Al + Si) ratio, with the lower Al content members of each group falling at the lower end of the ranges.

#### 5.3.4. Transparency Peak

[56] The 12-µm region of the reflectance spectra exhibits a reflectance maximum (transparency peak) due to enhanced volume scattering in fine grained materials, and occurs between the Si-O stretching (<12 µm) and bending (>12 µm) vibration bands (Figure 11) [Salisbury and Walter, 1989; Salisbury, 1993]. These investigators found that the wavelength position of this peak was negatively correlated with SCFM index. The peak is usually only apparent in unpacked powders containing some optically significant fraction of small (<~45 µm) grains. In the reflectance spectra, the wavelength position of this peak shows a positive correlation with Al:(Al + Si) ratio and a negative correlation with SCFM index (Figure 13). There is appreciably more scatter in the SCFM trend and the three data points which fall appreciably off the trend are all Narich samples (natrolite and phillipsite). These data suggest that this feature can be used to constrain Al:(Al + Si) ratio, in most cases generally to within  $\pm 5\%$ . As expected, there are no apparent correlations with major cation type or with structural group, since Si and Al contents are the major controls on the position of this feature.

#### 5.3.5. 12.5- to 15-µm Region

[57] Absorption features in the 12.5- to 15-µm range are commonly assigned to symmetric O-Si-O and O-Al-O stretching fundamentals [e.g., Moenke, 1974]. These bands are sensitive to Si-Al composition and shift to higher wavelengths with increasing Al content in transmission spectra [Breck, 1974]. The reflectance spectra of our samples show a diversity of spectral shapes in this region (Figure 14). This spectral region exhibits between one and three resolvable absorption bands. There is a general trend of increasing band position with increasing Al content but with significant scatter in the data. Many of the spectra exhibit overlapping absorption bands in this region, and using the first resolvable absorption band longward of 12 µm does not fully capture this spectral complexity. The sample spectra whose band positions fall off the general trend exhibit multiple overlapping absorption bands, and prominent inflections are present at the wavelength positions expected on the basis of their Al:(Al + Si) ratios. There are no apparent correlations between structural groups and either the number or relative strengths of absorption bands in this region, with the exception of the group 5 zeolites which exhibit the greatest number of resolvable absorption bands. As was the case with the absorption bands in the 8.5- to  $11-\mu m$  region, the position of the first major absorption band in the 12- to 15µm region does provide a means for discriminating some of the structural groups (Table 3) and the lower Al content members of each group fall at the lower end of the ranges. 5.3.6. 15.5- to 18-µm Region

[58] Absorption features in the 15.5- to 18-µm region should be present only in zeolites which contain double ring (D4R and D6R) structures, specifically groups 3 and 4; these features are assigned to O-Si-O and O-Al-O bends

Table 3. Wavelength Range of O-Si-O and O-Al-O Stretching and Bending Fundamentals by Structural Group

Structural Group		F	Range, μm		
	Ası	vmmetric Stret	ches		
1	8.8-9.3	9.7-10.3			
2	8.7	9.9 - 10.0			
4	8.6 - 8.8	10.0 - 10.1			
5	9.1-9.4	9.6-9.7	10.0 - 10.2	10.3-10.35	10.5-10.7
6	8.8	9.4-9.7			
7	8.5 - 8.8	9.9-10.3			
	Svi	mmetric Stretc	hes		
1	12.60-12.82				
2	12.60-12.63				
4	12.92-12.95				
5	12.50-14.12				
6	12.80				
7	12.50-12.70				
		Bends			
1	23.3 - 24.5				
2	22				
4	21.8 - 21.9				
5	21.3-22.1				
6	24.4-24.5				
7	23.8-24.9				



**Figure 13.** Wavelength position of the transparency peak in the 12- $\mu$ m region of the zeolite reflectance spectra as a function of composition: (a) versus Al:(Al + Si), (b) versus SCFM index.

[*Breck*, 1974]. Absorption bands in this region are present in a number of transmission and reflectance spectra of zeolites besides those from groups 3 and 4, but are most resolvable in the chabazite spectra (group 4). However, an absorption feature in this region is probably not diagnostic enough to permit its use in reliably discriminating type 3 and 4 zeolites.

#### 5.3.7. 20- to 26-µm Region

[59] Absorption features in the 20- to 26-µm region are largely assigned to O-Si-O and O-Al-O bending vibrations [*Stubičan and Roy*, 1961; *Breck*, 1974; *Moenke*, 1974; *Ryskin*, 1974]. Their wavelength positions are apparently not very sensitive to Si:Al ratios [*Breck*, 1974]. We found no well-defined correlations between wavelength positions of any of the bands in this region and Si:Al ratios. An added complication in searching for trends in this wavelength region is the lack of well-resolved absorption bands in many of the spectra (Figure 15). It was found, however, that the group 5 spectra consistently displayed the strongest absorption band in this region shortward of 22.2  $\mu$ m, while the remaining spectra exhibited the strongest absorption band longward of 22.2  $\mu$ m. These differences are probably related to the relative importance of particular bending vibrations for specific zeolite groups. In addition, the



**Figure 14.** 11- to 16- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites showing details of absorption features in this region. Some spectra have been vertically offset by the amount indicated in brackets for clarity. (a) Representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1; no offset); erionite (ZEO125; group 2; +0.05); chabazite (ZEO118; group 4; +0.1); natrolite (ZEO116; group 5; +0.13); mordenite (ZEO129; group 6; +0.25); stilbite (ZEO121; group 7; +0.3). (b) The same for group 5 zeolites; from bottom to top: thomsonite (ZEO109; no offset); scolecite (ZEO111; +0.06); mesolite (ZEO113; +0.15); natrolite (ZEO116; +0.2).



**Figure 15.** 20- to 26- $\mu$ m reflectance spectra of <45- $\mu$ m grain size fractions of zeolites. Some spectra have been vertically offset by the amount indicated in brackets for clarity. (a) Representative zeolites from each structural group. From bottom to top: gismondine (ZEO107; group 1; no offset); erionite (ZEO125; group 2; +0.05); chabazite (ZEO118; group 4; +0.12); natrolite (ZEO116; group 5; +0.16); mordenite (ZEO129; group 6; +0.25); stilbite (ZEO121; group 7; +0.3). (b) The same for group 5 zeolites; from bottom to top: thomsonite (ZEO109; no offset); scolecite (ZEO111; +0.06); mesolite (ZEO113; +0.15); natrolite (ZEO116; +0.16).

position of the first resolvable absorption band longward of 21  $\mu$ m allows some discrimination of the different structural groups (Table 3).

#### 6. Remote Sensing Implications

[60] Because water is generally not tightly bound in zeolites, has varying abundances, and water content and zeolite structure are often coupled [*Deer et al.*, 1963; *Delgass et al.*, 1969; *Gottardi and Galli*, 1985; *Tsitsishvili et al.*, 1992], zeolite spectra can be quite diverse, unlike most other silicate and aluminosilicate minerals. While the diversity of spectral shapes seen in the wavelength regions associated with H<sub>2</sub>O has some promise for analysis of pure phases, the situation is quite different from a remote sensing perspective. Given that zeolite spectra are dominated by H<sub>2</sub>O-, Si-, and Al-associated absorption features, their spectra are relatively simple [*Gaffney et al.*, 1984].

[61] The H<sub>2</sub>O-associated features can be easily obscured by other H<sub>2</sub>O/OH-bearing minerals and many of the metal-OH features in the 2.1- to 2.5- $\mu$ m region are also found in other mineral spectra [*Clark et al.*, 1990]. The high reflectance of zeolites in the visible and near-infrared wavelength region can easily be reduced, and the absorption bands reduced in intensity, by small quantities of intimately mixed opaque minerals [*Clark*, 1983; *Orenberg and Handy*, 1992]. In this sense they are similar to other common hydrated aluminosilicates such as kaolinite and montmorillonite [e.g., *Clark et al.* 1990].

[62] Other absorption features located below 3.3  $\mu$ m which might be useful for detection and discrimination of zeolites are associated with the O-H stretching and bending fundamentals, combination, and overtone bands, with the

exception of the Fe-associated features seen below 0.9  $\mu$ m. The main wavelength regions associated with water (1.4, 1.9, and 2.8  $\mu$ m) are areas where atmospheric water is strongly absorbing, leading to difficulties in obtaining high quality reflectance spectra with good signal-to-noise ratios and high spectral resolution necessary for zeolite detection and discrimination.

[63] In terms of remote sensing detection of zeolites on Mars, one factor which must be considered is that many zeolites undergo structural rearrangements during dehydration which may lead to spectral changes. Zeolites are expected to be readily susceptible to dehydration under current Martian surface conditions [Gooding, 1985; Ming and Gooding, 1988] and a high priority for our group for the future is to measure the spectra of zeolites under simulated Martian surface conditions [Craig et al., 2001]. This will allow us to assess how zeolite reflectance spectra react to exposure to Martian surface conditions (i.e., simulated temperatures, atmospheric pressure and composition, and ultraviolet light regime). More importantly we will also determine how the more diagnostic Si-O and Al-O bands in the 10-µm region are affected by this exposure, and whether these absorption bands are still useful for zeolite detection and discrimination.

#### 7. Conclusions

[64] The 0.3- to 26- $\mu$ m reflectance spectra of zeolites are characterized by high overall reflectance and all of the spectra exhibit a number of common absorption features such as H<sub>2</sub>O-associated absorption bands in the 1.4-, 1.9-, 2.7- to 3.1-, and 6.1- $\mu$ m regions, and Si- and Al-associated absorption bands in the 10- to 26- $\mu$ m region. The shapes and

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intensities of these bands are a function of a number of factors, some of which are intimately coupled. These include cation type, void size, and structural properties of the aluminosilicate framework. These in turn cause variations in the local electronic environment which affects properties such as degree of hydrogen bonding and interatomic distances. Such variations are expressed in the reflectance spectra as changes in the number, relative intensities, and wavelength positions of various absorption bands. The relationships between composition, structure, and spectral properties are sufficiently understood to permit discrimination of structural groups, individual zeolites, and identification of major cation types using reflectance spectra. However, in most geological settings, the presence of other minerals in zeolite-bearing assemblages will undoubtedly hinder zeolite detection and discrimination using reflectance spectroscopy.

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