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Abstract: In this work, Raman spectroscopic experiments are conducted on zeolites, including a total of 33 varieties and seven groups with different secondary structural frameworks, for which characteristic vibration modes are studied. Most of the zeolites show prominent Raman peaks in the spectral range between $200-1200 \text{ cm}^{-1}$. Different groups of zeolites can be recognized by differences in the wavenumbers of the T-O-T (T = Si and Al, O = oxygen) modes in the range $379-538 \text{ cm}^{-1}$, the M-O (M = metal,) modes in the range 250–360 cm^{-1} and the T-O bending modes in the range 530-575 cm⁻¹. All zeolites show characteristic Raman peaks in the range 379-529 cm⁻¹, except for natrolite group (fibrous) zeolites, which are characterized by T-O-T modes in the 433–447 cm⁻¹ range and T-O bending modes in the 528–538 $\rm cm^{-1}$ range. The analcime group (with singly connected four-ring chains) zeolites show T-O-T modes in the 379-392 cm⁻¹ and 475-497 cm⁻¹ ranges. The gismondine group (with doubly connected four-ring chains) zeolites have T-O-T modes in the $391-432 \text{ cm}^{-1}$ and $463-497 \text{ cm}^{-1}$ ranges. The chabazite group (with a six-cyclic ring) zeolites are characterized by M-O modes in the 320–340 cm⁻¹ range and T-O-T modes in the 477–509 cm⁻¹ range. The Raman modes of mordenite group zeolites (397–410 cm⁻¹ and 470–529 cm⁻¹) overlap with those of heulandite group zeolites ($402-416 \text{ cm}^{-1}$ and $480-500 \text{ cm}^{-1}$). Moreover, the mordenite group has a characteristic peak in the 502–529 cm⁻¹ range, and an additional peak in the 800–965 cm⁻¹ range. Another recognizable peak for the heulandite group is in the 612-620 cm⁻¹ range. The unknown zeolites (cowlesite) have unique characteristic peaks at 534 cm⁻¹, which can aid in the verification of their identity.

Keywords: zeolite; Raman; secondary structural framework

1. Introduction

Zeolites are commonly found minerals that are known for their important industrial applications. The term zeolite was proposed by Cronstedt in 1756, who found that these minerals expelled water when heated, as if they were boiling [1]. Dehydration and the absorption of moisture make them suitable for use as desiccants. The holes and vacancies in their relatively open structure allow the selective screening of ions of different sizes and can serve as a molecular sieve [2]. Previous researchers have studied the application of zeolites for the purification of livestock wastewater, landfill wastewater and aquaculture water [3,4]. The relationship between zeolite structures and the mechanisms of removing pollutants was further discussed in [5].



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). Surprisingly, the physical properties and geological importance of the zeolite group minerals were not well known until the late 20th century [6]. The formula for zeolites is more appropriately expressed as $(Li,Na,K)_a(Mg,Ca,Sr,Ba)_d[Al_{(a+2d)}Si_{n-(a+2d)}O_{2n}]\cdot mH_2O$ [6,7]. The letters a, d and n are considered for charges balanced with oxygen by the extraframe-work cations—usually Na, K, Ca, but less frequently Li, Mg, Sr, Ba (usually Si \geq Al and m \leq n). The basic structural unit of the crystal is the TO₄ tetrahedron, where T stands for Si and Al. TO₄ tetrahedrons are called primary building units [8]. Zeolite group minerals can be verified according to their differences in stacking and linking these tetrahedrons, which are called secondary framework units [9]. Based on this concept, Gottardi and Galli (1985) [6] classified zeolites into seven structural groups, the scheme of which was later revised by Ghobarkar et al. (2003) [10], as shown in Table 1.

Zeolite Group	Minerals	Zeolite Group	Minerals	
	Natrolite *		Gmelinite *	
-	Pranatrolite		Chabazite *	
-	Tetranatrolite		Willhendersonite	
Natrolite group	Mesolite	Chabazite group	Levyne *	
(fibrous)	Scolecite *	- (six-cyclic ring)	Erionite *	
	Gonnardite		Offretite	
	Edingtonite		Faujasite	
	Thomsonite *		Goosecreekite	
-	Analcime *		Gismondine *	
	Wairakite		Garronite *	
	Hsianghualite		Amicite *	
Analcimo groun	Viseite	- Cismondino group	Gobbinsite *	
(singly connected	Laumontite *	(doubly connected	Phillipsite *	
4-ring chains)	Leonhardite	four-ring)	Harmotome	
	Roggianite		Merlinoite *	
	Yugawaralite		Mazzite	
	Partheite	_	Paulingite	
	Pollucite	_		
	Mordenite *		Heulandite *	
-	Dachiardite		Clinoptilolite *	
Mordenite group	Epistilbite	Heulandite group	Stilbite *	
-	Ferrierite		Stellerite	
-	Bikitaite		Barrerite	
Unknown group	Cowlesite		Brewsterite	

Table 1. The classification of zeolite minerals and the samples adopted in this study [10]. Zeolite minerals used in this study are in bold type, and all of them were identified by X-ray powder diffraction (XRD) except pollucite and gonnardite.

* denotes zeolites that have been analyzed by energy dispersive spectroscopy (EDS) for chemical composition.

According to the scheme of Gottardi and Galli, 1985 and Ghobarkar et al., 2003 [6,10], more than 50 different species of zeolite have been identified, but only 47 natural zeolite minerals are classified into the seven groups: the natrolite group (fibrous zeolites), analcime group (singly connected four-ring chains), gismondine group (doubly connected four-ring chains), chabazite group (singly and doubly connected six-ring cyclic structure), mordenite group and heulandite group, in addition to cowlesite in the unknown structure group

(Table 1 and Figure 1). In this study, we focused on the investigation of Raman spectra of secondary building units of zeolites. Based on the results, we were also able to establish a Raman spectrum database.



Figure 1. Secondary framework units of zeolite groups: (**a**) fibrous structure, (**b**) singly connected four-ring chains, (**c**) doubly connected four-ring chains, (**d**) singly connected six-ring cyclic structure, (**e**) doubly connected six-cyclic structure, (**f**) mordenite structure and (**g**) heulandite structure (after Gottardi and Galli, 1985 [10]).

The phase identification of a mineral can be achieved by several methods. Conventional ways of identifying minerals by optical observation may present great difficulty when zeolite minerals are encountered. X-ray diffraction requires a sufficient quantity of the sample, which may not be readily available, and a powdered form needs to be prepared. The Raman spectroscopic method is a reliable tool for the non-destructive identification of mineral phases. It is easy to manipulate and does not require the tedious pre-treatment of a sample. Mineral phases can be identified quickly, provided that a complete set of Raman data for minerals are available. The RRUFFTM database is created by a RRUFF project from University of Arizona. They complete high quality spectral data from well characterized minerals and is developing the technology to share this information with the world. This database has provided Raman spectroscopic patterns for more than 4000 minerals. However, not all minerals are included; for example, zeolite minerals are only partially available.

In the past, Hong (2005) [11] and Chiang (2008) [12] reported Raman spectroscopic investigations on several zeolites. They attempted to perform a systematic comparison to test if the T-O-T modes could be adopted as a basis for the classification of zeolites and, furthermore, to find out if there exist any correlations in zeolites in their secondary framework units [6,10]. In their studies, some of the zeolite groups (such as the chabazite group in Hong's study and the mordenite group in Chiang's study) did not have sufficient samples to lead to a conclusive result. Apparently, more work needs to be done to justify their achievement. Despite the fact that valuable data can be obtained from the Internet, it is necessary for a systematic investigation of zeolites to be performed in order to obtain more meaningful data.

2. Method

Thirty-three zeolite species including natural and synthetic sources were used in this study. Zeolites that are rarely found in nature were synthesized by Chen (2002) [13,14]; for example gobbinsite, garronite, amicite and merlinoite. Almost all species were identified structurally by X-ray powder diffraction (XRD) or chemically by energy dispersive spectroscopy (EDS). XRD used D2 PHASER type from the Bruker Company in Karlsruhe, Germany. EDS was performed using an INCA-300 type from the Oxford Instruments in Buckinghamshire, United Kingdom, addition to the SEM of a JSM-6360LV type made by JEOL Company from Tokyo, Japan. The operating conditions were an acceleration voltage of 12–15 kV and an electric current of 0.18 nA, and every sample was analyzed at more than four spots. Each mineral was mounted in epoxy resin, and the surface was polished to a 5 μ m grain size. We chose albite and silica-alumina glasses for the standard.

Micro-Raman spectroscopy using a Labram HR VIS from the Horiba Jobin-Yvon Company (from Paris, France) was adopted for spectroscopic investigation. The laser source was a 532 nm green light diode laser. The magnification was 50X and the beam size was focused to a minimum diameter of about 2 μ m in a confocal manner. Before the experiment, the spectrometer was calibrated against the characteristic Raman peak of a silicon chip at 520 cm⁻¹ to be within an error of ± 1 cm⁻¹. Spectral ranges of 150–1200 cm⁻¹ were selected for the analysis and the data were processed by LabSpec 5 software. In each sample, three different spots were examined for better statistical results. The average time for each data acquisition was about 10 s.

3. Results and Discussion

3.1. The Raman Spectroscopy of Zeolites

The Raman spectroscopic results for each zeolite structure group are described as follows. In our results, the following should be noted:

- A. The numbers (1–3) before each zeolite mineral indicate the number of specimens of the same species, since each species may have had more than one specimen available for analysis.
- B. We compared our results with the RRUFF database and chose the data that matched best with our data in the tabulated lists.
- C. The stoichiometric composition and sample locality of each sample are listed.

3.1.1. Natrolite Group

In this study, the fibrous zeolite specimens (Figure 1a) included three thomsonite, one edingtonite, one scolecite and two mesolite samples and one gonnardite (Table 2) sample. In combination with the RRUFF data, these specimens show common characteristic peaks at 433–447 cm⁻¹ and 528–538 cm⁻¹ (Figure 2). In addition, there was another characteristic peak at 990–1100 cm⁻¹.

Our Raman data for natrolite are consistent with those reported by Goryainov (2001) [15]. Our data also agree well with the natrolite group reported by Wopeka (1998) [16].

3.1.2. Analcime Group

There are only three species for singly connected four-ring-chain zeolites (Figure 1b): analcime, laumontite and pollucite (Table 3). The numbers of this group are lower than the other zeolite groups. There were two specimens for analcime. Pollucite is a Cs-containing zeolite—a recently discovered mineral in this group. In combination with the RRUFF data, these specimens showed common characteristic peaks at 297–310 cm⁻¹, 379–392 cm⁻¹ and 475–497 cm⁻¹ (Figure 3). Only laumontite showed additional peaks at other ranges of the spectrum.

Zeolite	RRUFF No.	Composition	Locality
Thomsonite (3)	R050103	NaCa ₂ (Si ₅ Al ₅)O ₂₀ ·6H ₂ O Na _{0.76} Ca _{2.19} (Si _{4.96} Al ₅)O ₂₀ ·nH ₂ O	Folknare, North Bohemia, Czech Republic
Natrolite (2)	R040112	$\begin{array}{c} Na_{2}(Si_{3}Al_{2})O_{10}{\cdot}2H_{2}O\\ Na_{2.02}Ca_{0.01}(Si_{3}Al_{1.99})O_{10}{\cdot}nH_{2}O\end{array}$	Partridge Island, NS, Canada
Edingtonite	R040110	$Ba(Si_3Al_2)O_{10}\cdot 4H_2O$	Ice River, BC, Canada
Scolecite	R040111	Ca(Si ₃ Al ₂)O ₁₀ ·3H ₂ O Ca _{0.98} Si _{3.04} Al _{1.96} O ₁₀ ·nH ₂ O	Nashik, Maharashtra, India
Mesolite (2)	R050013	Na ₂ Ca ₂ (Si ₉ Al ₆)O ₃₀ ·8H ₂ O	Aurangabad, Maharashtra, India
Gonnardite		$(Na, Ca_{0.5})_{8-10}(Al_{8+x}Si_{12-x})O_{40} \cdot 12H_2O_x = 0-2$	Schellkopf, Brenk, Niederzissen, Eifel, Rhineland-Palatinate, Germany

 Table 2. Natrolite group zeolite specimens (chemical formula in blue analyzed by EDS).



Figure 2. Raman spectra of natrolite group zeolites.

No.	RRUFF No.	Composition	Locality
Analcime (2)	R040128	NaSi ₂ AlO ₆ ·H ₂ O Na _{0.95} Si _{2.04} Al _{0.96} ·nH ₂ O	Wasson's bluff, Bay of Fundy, NS, Canada
Laumontite	R040020	$\begin{array}{c} Ca(Si_{4}Al_{2})O_{12}{\cdot}4H_{2}O\\ Na_{0.02}Ca_{1}(Si_{4.01}Al_{1.98})O_{12}{\cdot}nH_{2}O \end{array}$	Ozar, Nashik, India
Pollucite	R050344	$Cs(Si_2Al)O_6 \cdot nH_2O$	Chamachho, Skardu, Pakistan

Table 3. Analcime group zeolite specimens (chemical formula in blue analyzed by EDS).



Figure 3. Raman spectra of analcime group zeolites.

3.1.3. Gismondine Group

There are more mineral species in the doubly connected four-ring-chain zeolite group (Figure 1c), including specimens of harmotome, paulingite, gismondine, garronite, amicite, merlinoite and gobbinsite, in addition to two phillipsite specimens (Table 4). Among them, the rarely seen natural minerals of gismondine, garronite, amicite, merlinoite and gobbinsite are synthesized from glass through hydrothermal processes [13,14]. All have corresponding data in the RRUFF files except for gobbinsite. In combination with the RRUFF data, these specimens showed common characteristic peaks at 391–432 cm⁻¹ and 463–497 cm^{-1} (Figure 4). The most intense peak in this group, spanning the range 463–497 cm⁻¹, was found to be due to the relatively low wavenumber in gobbinsite and relatively high wavenumber in paulingite compared to the other members in this group. Otherwise, the most intense peak spanned a smaller range of 467–486 cm⁻¹. Gismondine, as reported by Mozgawa (2001) [17], had its most intense peak at 1086 cm^{-1} , which is not reasonable. Another reason for the wide span of characteristic peaks is the difference in the Si/Al ratio. In the process of synthesis, gismondine and amicite have lower Si/Al (~1), while gobbinsite and garronite have higher Si/Al (\sim 1.5–2.2) [13]. This may influence the location of the highest peak between 463-497 cm⁻¹.

3.1.4. Chabazite Group

We analyzed one specimen of each of the six-ring cyclic structure zeolites (Figs. 1d and 1e), including chabazite, levyne, goosecreekite, gmelinite and erionite (Table 5). In combination with the RRUFF data, these specimens showed common characteristic peaks at 320–340 cm⁻¹ and 477–509 cm⁻¹ (Figure 5). In addition, chabazite, levyne and erionite had peaks at 460 cm⁻¹, 466 cm⁻¹ and 470 cm⁻¹, respectively. Among them, only goosecreekite, which was recently found and classified as a member of the chabazite group, lacked the peak at 320–340 cm⁻¹. There are no results for goosecreekite in the RRUFF data bank. We

suspect that goosecreekite may not belong to the chabazite group. Further investigation is needed to verify the identity of goosecreekite.

Table 4. Gismondine group zeolite specimens (chemical formula in blue analyzed by EDS).

No.	RRUFF No.	Composition	Locality
Harmotome	R070015	$Ba_2(Si_{12}Al_4)O_{32}\cdot 12H_2O$	Strontian, Argylshire, Scotland, UK
Paulingite	R070604 R060457	(K,Ca,Na,Ba) ₁₀ (Si,Al) ₄₂ O ₈₄ ·34H ₂ O (Na ₂ ,K ₂ ,Ca,Ba) ₅ Al ₁₀ Si ₃₅ O ₉₀ ·45H ₂ O	Vinařice, Kladno, Central Bohemia, Czech Republic
Gismondine	R060809	Ca ₂ (Si ₄ Al ₄)O ₁₆ ·8H ₂ O (Na _{0.05} K _{0.01} Ca _{2.17})(Si _{3.9} Al ₄)O ₁₆ ·nH ₂ O	Arensberg, Zilsdorf, Walsdorf, Gerolstein, Vulkaneifel District, Rhineland-Palatinate, Germany
Garronite	R050281	$\begin{array}{c} Ca_3(Si_{10}Al_6)O_{32}{\cdot}14H_2O\\ (Na_{0.08}K_{0.02}Ca_{2.53})(S_{10.65}A_{5.42})O_{32}{\cdot}nH_2O \end{array}$	Synthetic zeolite from Chen et al. (2002) [13]
Amicite	R080066	$Na_{2}K_{2}(Si_{4}Al_{4})O_{16}\cdot 5H_{2}O \\ Na_{1.52}K_{1.52}Ca_{0.07}Si_{4.29}Al_{3.89}O_{16}\cdot nH_{2}O$	Synthetic zeolite from Chen et al. (2002) [13]
Merlinoite	R130095	$\begin{array}{c} K_5 Ca_2(Si_{23}Al_9)O_{64}{\cdot}24H_2O \\ (Na_{2.14}K_{7.06}Ca_{0.19})(Si_{22.06}Al_{10.05}){\cdot}nH_2O \end{array}$	Synthetic zeolite from Chen et al. (2002) [14]
Gobbinsite	n/a	$\begin{array}{l} Na_{5}(Si_{11}Al_{5})O_{32}\cdot 11H_{2}O\\ (Na_{5.08}K_{0.01}Ca_{0.02})(Si_{10.73}Al_{5.32})O_{32}\cdot nH_{2}O\end{array}$	Synthetic zeolite from Chen et al. (2002) [13]
Phillipsite 1, 2	R050078	$\hline \hline \hline Ca_3(Si_{10}Al_6)O_{32}\cdot 12H_2O \\ (K_{1.34}Na_{0.08}Ca_{1.2})(Si_{12.19}Al_{3.81})\cdot nH_2O$	Rydec, Usti nad Labem, North Bohemia,
	R070271	$Na_6(Si_{10}Al_6)O_{32}\cdot 12H_2O$	Czech Republic

Table 5. Chabazite group zeolite specimens (chemical formula in blue analyzed by EDS).

No.	RRUFF No.	Composition	Locality
	R050014	$Ca_2Si_8Al_4O_{24}\cdot 13H_2O$	
Chabazite	R120032	(K ₂ NaCa _{0.5})Si ₈ Al ₄ O ₂₄ ·11H ₂ O K _{2.70} Na _{0.02} Ca _{0.99} Si _{8.68} Al _{2.86} O ₂₄ ·nH ₂ O	Wasson's bluff, bay of Fundy, Novascotia, Canada
	R061095	$(Na_3K)Si_8Al_4O_{24}\cdot 11H_2O$	
Levyne	R070270	$\begin{array}{c} Ca_{3}(Si_{12}Al_{6})O_{36}{\cdot}18H_{2}O\\ (Na_{1.29}K_{0.26}Ca_{1.8})(Si_{12.19}Al_{6.04}){\cdot}nH_{2}O\end{array}$	Moonen Bay, Isle of Skye, Scotland, UK
	R060021	$Na_6(Si_{12}Al_6)O_{36}{\cdot}18H_2O$	
Goosecreekite	n/a	$CaSi_{6}Al_{2}O_{16}\cdot 5H_{2}O$	Nashik, Maharashtra, India
Gmelinite	R060168	$\begin{array}{l} Na_4(Si_8Al_4)O_{24}{\cdot}11H_2O\\ (Na_{2.19}K_{1.21}Ca_{0.25})(Si_{8.12}Al_{3.87})O_{24}{\cdot}nH_2O \end{array}$	Five Island, NS, Canada
Erionite	R060836	$\begin{array}{c} Ca_5(Si_{26}Al_{10})O_{72}\cdot 30H_2O\\ (Ca_{1.76}Mg_{0.71}Na_{0.47}K_{2.58})\\ (Si_{28.19}Al_{7.76})O_{72}\cdot nH_2O \end{array}$	Little Ajo Mountains, AZ, USA

3.1.5. Mordenite Group

Zeolites of the mordenite group (Figure 1f) are also rare; therefore, only one specimen from each of epistilbite, mordenite, dachiardite and bikitaite was analyzed (Table 6). The Raman spectra of zeolites in the mordenite group were more diversified (Figure 6). Knops-Gerrits et al. (1997) [18] reported that mordenite has its most intense peak at 395 cm⁻¹, which is consistent with our mordenite [18]. According to Monsgawa (2001) [17], the most intense peak of mordenite is at 534 cm⁻¹, which is different from our observation and the RRUFF data. In addition, the dachiardite of Mozgawa (2001) [17] had its most intense peak at 714 cm⁻¹, which is also different from our result and the RRUFF data on dachiardite.



We suspect that the result of Mozgawa (2001) [17] for dachiadite may not be reliable since most of the zeolites have their most intense peak in the spectral range of 379-538 cm⁻¹.

Figure 4. Raman spectra of gismondine group zeolites.

No.	RRUFF No.	Composition	Locality
Epistilbite	R061105	$Ca_3(Si_{18}Al_6)O_{48}\cdot 16H_2O$	Ojhar mine, Nashik, Maharashtra, India
Mordenite	R061118	$\begin{array}{c}(\mathrm{Na}_2,\mathrm{Ca},\mathrm{K}_2)_4(\mathrm{Si}_{40}\mathrm{Al}_8)\mathrm{O}_{96}\cdot28\mathrm{H}_2\mathrm{O}\\(\mathrm{Na}_{1.24}\mathrm{K}_{1.45}\mathrm{Mg}_{0.03}\mathrm{Ca}_{2.05})\\(\mathrm{Si}_{40.93}\mathrm{Al}_{7.14})\mathrm{O}_{96}\cdot\mathrm{nH}_2\mathrm{O}\end{array}$	Nashik, Maharashtra, India
Dachiardite	R061097	Ca ₂ (Si ₂₀ Al ₄)O ₄₈ ·13H ₂ O	Pahlia river, Culverden, Canterbury, South Island, New Zealand
Bikitaite	R050160	$Li(AlSi_2O_6) \cdot H_2O$	Kings Mountain, NC, USA

Table 6. Mordenite group zeolite specimens (chemical formula in blue analyzed by EDS).



Figure 5. Raman spectra of chabazite group zeolites.



Figure 6. Raman spectra of mordenite group zeolites.

The epistilbite in the RRUFF data bank (R061105) shows peaks at 397, 438, 469 and 805 cm⁻¹, but our epistilbite only showed peaks at 410 and 438 cm⁻¹. However, our Raman spectrum for epstilbite agrees well with that of Mozgawa (2001) [17]. It is likely that R061105 may contain Raman signals from two minerals, with most of its characteristic

peaks similar to those of mordenite. However, our epistilbite specimen is a single crystal, which yields a more reliable Raman spectrum. In short, in combination with the RRUFF data, these specimens showed more than one characteristic peak in the spectral range $397-470 \text{ cm}^{-1}$, and another peak in the spectral range $502-529 \text{ cm}^{-1}$. In this group, an additional peak appeared in the spectral range of $800-965 \text{ cm}^{-1}$.

3.1.6. Heulandite Group

We had a complete set of zeolites in the heulandite group (Figure 1g), including two specimens of clinoptilolite, stellerite and stilbite, and one specimen of brewsterite, stilbite, barrerite and heulandite (Table 7). In combination with the RRUFF data, these specimens showed common characteristic peaks at 402–416 cm⁻¹, 480–500 cm⁻¹ and 612–620 cm⁻¹ (Figure 7).



Figure 7. Raman spectra of heulandite group zeolites.

3.1.7. Unknown Structure

Cowlesite is not classified as belonging to any of the zeolite structural groups yet (Table 8). The Raman spectrum of cowlesite was not very complicated and only showed peaks at 325, 393 and 534 cm⁻¹. The peaks at 325 and 393 cm⁻¹ were not very prominent (Figure 8). Since no other zeolites are classified in this group, we tentatively take 534 cm⁻¹ as the characteristic peak for zeolites of the unknown group.

No.	RRUFF No.	Composition	Locality	
Provestorito	R070120	$Ba(Al_2Si_6)O_{16}{\cdot}5H_2O$	Yellow Lake, Osoyoos Mining Division,	
R070227		$Sr(Al_2Si_6)O_{16} \cdot 5H_2O$	BC, Canada	
$\frac{Ca_{3}(Si_{30}Al_{6})O_{72}\cdot 20H_{2}O}{(Ca_{1.7}Mg_{0.03}Na_{1.31}K_{1.07})(Si_{29.69}Al_{6.47})O_{72}\cdot nH_{2}O}$				
Clinoptilolite (2)	R110201	$K_6(Si_{30}Al_6)O_{72} \cdot 20H_2O$	Malheur co., OR, USA	
	R061099	$Na_6(Si_{30}Al_6)O_{72}\cdot 20H_2O$		
Stellerite (2)	R040174	$Ca_4(Si_{28}Al_8)O_{72}\cdot 28H_2O$	Tisgaon Mine, Aurangabad, Maharashtra, India	
Stilbite (2)	R050012	NaCa4(Si ₂₇ Al9)O ₇₂ ·28H ₂ O (Na _{0.46} K _{0.01} Ca _{4.06})(Si _{27.79} Al _{8.08})O ₇₂ ·nH ₂ O	Nashik, Maharashtra, India	
Barrerite	R050135	$Na_2(Al_2Si_7)O_{18}\cdot 6H_2O$	Alaska garnet Mines, Kuiu Island, AK, USA	
	R050017	NaCa4(Si27Al9)O72·24H2O (Na1.89K0.15Ca3.26)(Si27.69Al8.23)O72·nH2O		
Heulandite	R061137	KCa4(Si27Al9)O72·24H2O	Nashik, Maharashtra, India	
	R070281	(Na,Ca) ₆ (Si,Al) ₃₆ O ₇₂ ·24H ₂ O		
-	R070272	$NaSr_4(Si_{27}Al_9)O_{72}\cdot 24H_2O$		

Table 7. Heulandite group zeolite specimens (chemical formula in blue analyzed by EDS).

Table 8. Unknown structure group zeolite specimen.

No.	RRUFF No.	Composition	Locality
Cowlesite	R060627	$Ca(Al_2Si_3)O_{10}{\cdot}5{-}6H_2O$	County Antrim, Northern Ireland, UK



Figure 8. Raman spectra of cowlesite.

3.2. Comparison between Different Zeolite Groups

According to Gujar et al. (2005) [19], the Raman peaks in the spectral range of $380-530 \text{ cm}^{-1}$ represent the vibration modes of T-O-T bonding, where T stands for Si or Al. In this study, we have integrated our results for Raman spectra in Table 9. All zeolites invariantly showed two separated regions of peak locations in the spectral range of $379-538 \text{ cm}^{-1}$. One location was in the spectral range $379-447 \text{ cm}^{-1}$ and the other was in the spectral range of $463-538 \text{ cm}^{-1}$. We tentatively suggest that the former and the latter sets of peaks may be associated with Al-O-Al and Si-O-Si, respectively. All the Al-O-Al modes were in the spectral range of $379-447 \text{ cm}^{-1}$ except for the chabazite group. However, zeolites in the chabazite group showed another characteristic peak at $320-340 \text{ cm}^{-1}$, which can be used as a criterion to distinguish it from other groups. The relatively complicated structure of the mordenite and heulandite groups showed additional peaks in the spectral ranges of $612-620 \text{ cm}^{-1}$ and $800-965 \text{ cm}^{-1}$, which were also unique among the zeolite groups. Another interesting phenomenon was the most intense peak at $463-538 \text{ cm}^{-1}$ for

most zeolite groups except the mordenite and heulandite groups, which showed their most intense peak at 397-416 cm⁻¹.

Al-O-Al Modes	Si-O-Si Modes	Others
433–447	528-538	
379-392	475-497	
391-432	463-497	
	477-509	320-340
397-410	470-529	800–965
402–416	480-500	612–620
	Al-O-Al Modes 433–447 379–392 391–432 397–410 402–416	Al-O-Al Si-O-Si Modes Modes 433-447 528-538 379-392 475-497 391-432 463-497 477-509 397-410 470-529 402-416

Table 9. The Al-O-Al and Si-O-Si modes of each group of zeolites (unit: cm^{-1}).

After carefully comparing the characteristic Raman peaks of all zeolites, we obtained the following rules for the identification of different zeolite groups, as shown in Figure 9.

Zeolite group		Raman shift (cm ⁻¹)						
Natrolite group	350	400 4	50 500	8-538	600	700	22	1000
Analcime group		379-392	475-497				~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
Gismondine group		391-432	463-497				~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
Chabazite group	320-340		477-509					
Mordenite group		397-410	470-529				800-9	65
Heulandite group		402-416	48 <mark>0-5</mark> 00		612 <mark>-</mark> 620		22	

Figure 9. Characteristic Raman spectrum peaks of each group of zeolites.

- (1) The fibrous natrolite group can be identified by the characteristic peaks at 528-538 cm⁻¹.
- (2) The singly-connected four-cycle chain analcime group can be identified by the characteristic peaks at 379–392 cm⁻¹.
- (3) The doubly-connected four-cycle chain gismondine group has a pattern almost overlapping the zeolites of other groups, and only the characteristic peaks at 391–432 cm⁻¹ can be used for the recognition of this group. The Al-O-Al modes of gismondine, amicite and gobbinsite fall in the spectral range of 391–432 cm⁻¹, which overlaps with the Al-O-Al modes in the mordenite and heulandite groups. This group has a very wide composition.
- (4) The six-cycle ring chabazite group can be recognized by the unique presence of a characteristic peak at 320–340 cm⁻¹.
- (5) The spectra of the mordenite group overlap with other zeolites, and the peaks at 800-965 cm⁻¹ are suitable for its identification.
- (6) The spectra of the heulandite group also overlap with other zeolites in prominent characteristic peaks, and only the peak at 612–620 cm⁻¹ can be used for its identification.
- (7) Most mordenite and heulandite groups have higher intense peaks in the range of Al-O-Al modes, which are different from the gismondine group.

3.3. The Meaning of Characteristic Vibration Modes

Zeolite minerals are numerous, and their structures are diverse. Therefore, it is difficult to distinguish their identity, irrespective of which XRD or Raman spectroscopic method is applied for the identification. It is quite common for more than two zeolite minerals to coexist in a rock due to their similarity in occurrence. The presence of solid solution in zeolites [13,14,20] may also cause a shift in the position of Raman characteristic peaks [11,21,22]. The results of this study provide the first-principle qualitative identification of zeolites among various structural groups, which makes the further classification of zeolites easier.

Mozgawa (2001) [17] analyzed the vibrational modes of zeolites by means of IR and Raman spectroscopy and concluded that the 390–415 cm⁻¹ and 480–500 cm⁻¹ ranges are

associated with five-ring (heulandite group) and four-ring zeolites, respectively. This structural relationship was confirmed by Yu et al. (2001) [23], who concluded that 370–430 cm⁻¹ and 470–530 cm⁻¹ modes are related to five-ring and four-ring zeolites, respectively [23]. Yu et al. (2001) [23] further proposed that peaks at 290–410 cm⁻¹ and 220–280 cm⁻¹ can be related to six-ring and eight-ring zeolite structures, respectively.

Mozgawa (2001) [17] also concluded that IR provides more significant evidence than Raman spectra regarding the relationship of vibrational modes and structural types in zeolites, which is currently under investigation. However, we believe that our systematic work on zeolites indicates that Raman spectroscopic data can also provide sufficient evidence for the assignment of vibrational modes to structural types in zeolite.

The physical significance of various vibrational modes was investigated by Gujar et al. (2005) [19] who compiled previous studies on zeolites and assigned their vibration modes as shown in Table 10. These modes are mostly attributed to the linear stretching vibration and non-linear bending vibration modes. All zeolite minerals have their characteristic peaks in the spectral range of 380–538 cm⁻¹, which denotes the T-O-T vibration modes. As described above, these peaks have been tentatively assigned as the Al-O-Al and Si-O-Si modes. The relative intensity and wavenumber of the Al-O-Al and Si-O-Si modes may be influenced by the Al/Si, which is an issue that needs further study. As for the M-O vibration mode, only the chabazite group shows obvious peaks at 320–340 cm⁻¹. The characteristic peaks in the range of 800–965 cm⁻¹ may be due to the vibration of Si in the tetrahedrons, which only occurs in more complex structures such as the mordenite and heulandite groups. Further investigation regarding zeolites of new species or species with unknown structures is required.

Wavenumber Range (cm $^{-1}$)	Vibration Mode
250–360	M-O (M stands for cations)
379–447	O vibration in T-O-T (T stands for Al)
459–538	O vibration in T-O-T (T stands for Si)
530–575	T-O bending
780–980	Si in tetrahedron
1100–1450	T-O bending
3224, 3474	OH stretching
3550	OH stretching (bridging)

Table 10. Raman vibration modes of zeolites (modified from Ref. [19]).

4. Conclusions

The most intense Raman peaks occur in the spectral range of 379–538 cm⁻¹, representing the T-O-T vibrational modes, almost invariantly containing the Al-O-Al and Si-O-Si modes. The fibrous natrolite group is most easily recognized by the specific T-O-T vibrational modes at 433–447 cm⁻¹ and the T-O mode at 528–538 cm⁻¹. The singly-connected four-ring chain analcime group processes a specific Al-O-Al mode at 379–392 cm⁻¹, which is separable from other groups of zeolites. The six-ring chabazite group lacks an Al-O-Al mode, but the specific M-O mode at 320–340 cm⁻¹ is unique among all zeolites. In the mordenite group, additional modes of Si-O-Si at 470–529 cm⁻¹ and Si in the tetrahedron mode at 800–965 cm⁻¹ can be used as a criterion to distinguish it from other groups. The heulandite group stands apart due to the presence of a vibrational mode at 612–620 cm⁻¹. The doubly-connected four-ring chain of the gismondine group overlaps its T-O-T mode with the mordenite and heulandite groups, but the gismondine group has minor T-O-T modes at 391–432 cm⁻¹. In the gismondine group, only gobbinsite has no specific criterion for the recognition of its identity, which is an issue that requires further research or more detailed spectral investigation.

Band positions correspond only to the samples tested in this study. The band positions may differ due to different types of equipment, excitation laser source, chemical composition and crystal orientation when results from other laboratories are compared. More work needs to be done to justify the achievements in this study. Despite the fact that numerous data can be obtained through websites such as RRUFF, it is necessary that systematic investigations of zeolites are performed in order to obtain meaningful results.

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Data Availability Statement: The data presented in this study are available on request from the corresponding author Huei-Fen Chen, because most data were established in our lab. The data are not publicly available due to "we want to construct a software for mineral identification by Raman spectrum peaking" in the future.

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