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THE JOURNAL OF PHYSICAL CHEMISTRY A-

Raman and Infrared Spectroscopy of α and β Phases of Thin Nickel Hydroxide Films Electrochemically Formed on Nickel

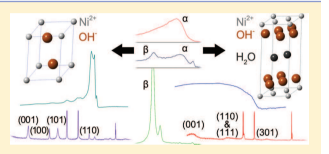
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ABSTRACT: The present work utilizes Raman and infrared (IR) spectroscopy, supported by X-ray diffraction (XRD) and X-ray photoelectron spectroscopy (XPS) to re-examine the fine structural details of Ni(OH)₂, which is a key material in many energy-related applications. This work also unifies the large body of literature on the topic. Samples were prepared by the galvanostatic basification of nickel salts and by aging the deposits in hot KOH solutions. A simplified model is presented consisting of two fundamental phases (α and β) of Ni(OH)₂ and a range of possible structural disorder arising from factors such as impurities, hydration, and crystal defects.



For the first time, all of the lattice modes of β -Ni(OH)₂ have been identified and assigned using factor group analysis. Ni(OH)₂ films can be rapidly identified in pure and mixed samples using Raman or IR spectroscopy by measuring their strong O–H stretching modes, which act as fingerprints. Thus, this work establishes methods to measure the phase, or phases, and disorder at a Ni(OH)₂ sample surface and to correlate desired chemical properties to their structural origins.

1. INTRODUCTION

The oxides and hydroxides of Ni have been extensively studied for many potential applications including energy storage,¹⁻¹² electrochromic devices,^{13,14} and photocatalysis.¹⁵ Corrosion studies of Ni have focused on the investigation of the nature of its surface layers and passivation phenomena.^{16–19}

Two phases of Ni(OH)₂, denoted α and β , were first established from X-ray diffraction (XRD) patterns,² and their physical properties have since been thoroughly investigated utilizing infrared spectroscopy (IR),^{1,4,6,7,14,20–28} Raman spec-troscopy,^{5,22,29–39} extended X-ray absorption fine structure (EXAFS),^{30,40} inelastic neutron scattering (INS),⁴¹ neutron diffraction,^{42–44} and more detailed XRD studies.^{10,31,45,46} Typically, Ni(OH)₂ is prepared by the basification of an aqueous nickel(II) salt solution either by the addition of an alkali hydroxide or ammonia or by the electrochemical reduction of water. The joint effects of current density or base solution concentration, nickel concentration, and the counterions determine the product phase, crystallinity, and, in the electrochemical approach, the current efficiency.47,48 Literature references to Ni(OH)₂ that do not specify the phase usually refer to well-crystallized, high purity β -Ni(OH)₂. β -Ni(OH)₂ is available commercially. This phase, which occurs in nature as theophrastite, has hexagonal scalenohedral symmetry and is isostructural to brucite $[Mg(OH)_2]^{2,11,49-51}$ Most generally, α -Ni(OH)₂ has trigonal symmetry consisting of planes of β -Ni(OH)₂ intercalated with water. This hydroxide has been given the general formula Ni(OH)_{2-x} $A_{x/n}$ ·yH₂O, where $0 \le x \le 1, 0$ $\leq y \leq 2$, and *n* is the anionic charge of an impurity A^{n-2}

However, the existence of intermediate phases due to disorder, hydration and the incorporation of impurities has been proposed and numerous methods of denoting these materials exist, including $\alpha_{\rm am}$, α^* , α' , IS, and $\beta_{\rm bc}$.^{11,21,52,53} These notations are used inconsistently between reports by different authors and are often ambiguous and redundant.

Several studies have linked structural changes in Ni(OH), electrodes and devices to their performance. As an example, the Ni(OH)₂/NiOOH redox pair is used extensively for the anodes of secondary alkaline batteries, including NiCd and NiMH,⁵⁴ and it is generally agreed to be limited by electron and proton conductivities, which depend on the crystal microstructure.⁴¹ In one study, the incorporation of Co impurities in a β -Ni(OH)₂ electrode was observed to improve its charge capacity, although the simultaneous incorporation of other types of structural disorder was not considered, and therefore, it cannot be conclusively stated whether structural disorder or impurities are responsible for the performance improvements.²⁹ In another example, the electrochromic efficiency and stability during electrochemical cycling of precipitated Ni(OH)₂ films were linked to the nickel salts used during preparation, although, without further details of the structural differences, the authors were unable to determine the origin of the differences between samples.¹⁴ This demonstrates the need for methods to measure the incorporation of anions from commonly used nickel salts,

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such as sulfate, chloride, and nitrate, and separately examine other types of disorder, such as hydration and crystal defects. Such work will allow more systematic investigations of the link between structural changes and material performance.

Although impurities and defects can improve electrode performance, they can also cause instability; for instance, α -Ni(OH)₂ is not currently used in battery applications because its charge capacity tends to decrease more rapidly on cycling than that of β -Ni(OH)₂ electrodes.²⁹ One can consider, as an example, that the slow leaching of impurities could cause other problems, e.g., chloride anions initiate pitting corrosion of certain metals that may be used elsewhere in a device. Thus, methods to clearly identify different types of disorder and impurities in the different phases of Ni(OH)₂ will allow useful disorder to be incorporated while simultaneously deliberately avoiding problematic disorder that destabilizes the material.

The introduction of such terms as active/deactivated α and active/deactivated β to describe Ni(OH)₂ electrodes demonstrates a lack of knowledge about both of these phases and reaffirms their practical importance.⁵² A more complete description of the two main phases of Ni(OH)₂ and any intermediates will be invaluable to research focused on applied Ni(OH)₂ electrochemistry. The ability to measure and, ultimately, to control the types of disorder present in an electrode or catalyst material will allow for the optimization of its performance, as determined by charge capacity, electrochromic efficiency, or catalytic activity, while simultaneously enabling improvements in the long term stability of an electrode or device.

The present work utilizes Raman and IR vibrational spectroscopy using modern instruments that offer higher resolution and sensitivity than previously reported, supported by XPS and fixed incidence XRD measurements, to re-examine the fine structural details of $Ni(OH)_2$. It is from the combination of these four analytical techniques with a review of previous studies that we now present a comprehensive, yet simplified, model of $Ni(OH)_2$ that considers its possible phases, incorporation of impurities and structural disorder. For example, our XRD patterns of β -Ni(OH)₂ show crystallographic disorder, which allows us to rationalize the appearance of additional, and previously unidentified, stretching modes in the Raman and IR spectra. As another example, N peaks in the XPS spectra of α -Ni(OH)₂ prepared from a Ni(NO₃)₂ solution indicates that nitrate anions are incorporated in its structure. By observing the relative intensities of Raman and IR peaks for the sample, we conclude that the nitrate anions occupy at least two distinct chemical environments. This study identifies and assigns, for the first time, all of the lattice modes of β -Ni(OH)₂ using factor group analysis and summarizes the known Raman and IR peaks for both α - and β -Ni(OH)₂.

2. EXPERIMENTAL METHODS

2.1. Material Preparation. Metallic Ni foil (Alfa Aesar, \geq 99%, 0.127 mm) electrodes (1 cm × 2 cm) were attached to Ni wire (Alfa Aesar, 99.9%, 0.065 mm diameter) by electric arc welding. The wire and one end of the electrode were covered with Teflon tape. The Ni foil substrates were washed with acetone and ethyl alcohol before sonicating them in high purity water (Millipore Milli-Q, 18.2 M Ω cm). Immediately after, the substrates were cleaned in an acid bath (50% acetic acid, 30% phosphoric acid, 10% sulfuric acid, 10% nitric acid, 85 °C) for 30 s then rinsed again with excessive amounts of high purity water.

Ni hydroxide films were formed by the conditions and treatments listed in Table 1. A Ni wire ($\sim 10 \text{ cm long}$) served as a

Article

Table 1. Methods Used for Sample Preparation

sample	start material	current density (mA cm ⁻²)	time	solution	temperature (°C)
α_1	substrate	-2.5	10 min	0.1 M NiSO4	22
α_2	substrate	-2.5	10 min	$0.1 \mathrm{~M~NiCl}_2$	22
α_3	substrate	-2.5	2 min	0.1 M Ni(NO ₃) ₂	22
β_1	α_1	N/A	185 h	30% KOH	90
β_2	α_2	N/A	185 h	30% KOH	90
β_3	α_3	N/A	185 h	30% KOH	90
lpha/eta	α_2	N/A	140 h	Water	22

counter electrode. Electrochemical treatments were performed using a Solartron Analytical 1470E multistat controlled with MultiStat software (v. 1.5a, Scribner Associates Inc.). Deposition solutions were prepared from NiSO₄·6H₂O (Alfa Aesar, \geq 98%), NiCl₂·6H₂O (J.T. Baker, assay = 100.7%), and Ni(NO₃)₂·6H₂O (Alfa Aesar, \geq 98%). All solutions were prepared with high purity water. Solution concentrations were selected for efficient precipitation based on reported studies.⁴⁸ The current density and deposition times were selected to form α -Ni(OH)₂ films generally sufficiently thick to be identified by the spectroscopic methods. Aging solutions were prepared from KOH pellets (Fisher Scientific, ACS certified).

Sample α_1 was prepared in duplicate, and each sample was analyzed by XRD, XPS, Raman spectroscopy, and FT-IR spectroscopy to confirm reproducibility in the sample preparation and measurement methods. Sample α_3 was also prepared twice, and each sample was analyzed by XPS, Raman, and FT-IR spectroscopy to further confirm reproducibility. Samples β_1 , β_2 , and β_3 were stored in high purity water after aging in base and were dried immediately before analysis. The preparation methods for the β -phase samples were considered to be wellcontrolled and reproducible because the α -phase samples were found to be reproducible, and the measurements for the three β -phase samples gave similar results.

2.2. X-ray Diffraction (XRD). Samples were analyzed using a Bruker AXS D8 DISCOVER diffractometer. Fixed incidence (5°) measurements were taken at 0.02° point⁻¹ and 3 s point⁻¹ using a Cu K α X-ray source ($\lambda = 1.54$ Å). Data were smoothed for presentation with a 10-point, second order Savitzky–Golay filter.

2.3. X-ray Photoelectron Spectroscopy (XPS). Samples were analyzed with a Kratos Analytical Axis Ultra DLD equipped with a mono Al $K\alpha$ X-ray source. Survey spectra, used for elemental quantification, were collected at 160 eV pass energy, and high-resolution spectra of the Ni 2p and the C 1s regions were collected at 20 eV pass energy. All spectra were collected at a 54° takeoff angle. Spectra were processed with CasaXPS (v. 2.3.16dev95, Casa Software Ltd.) using Shirley backgrounds and the Ni 2p peak positions and widths for β -Ni(OH)₂, NiO, and Ni metal reported by Biesinger et al. The elemental relative sensitivity factors (RSF) provided by Kratos Analytical were used for elemental quantification. Measured binding energies were adjusted to compensate for the shift that arises from a buildup of electrostatic charge by setting the adventitious hydrocarbon C 1s peak to the binding energy 284.8 eV.⁵⁵

2.4. Raman Spectroscopy. The Raman measurements were performed at room temperature ($22 \degree C$) employing 1 mW of 457 nm laser light (Cobalt Twist diode-pumped solid state laser) for excitation in a backscattering geometry using the confocal microscope with a 100× objective on a Jobin-Yvon T64000 triple

spectrometer operated in the subtractive mode and equipped with a back-illuminated Si charge-coupled-device detector. The sample surfaces, in general, were not planar, comprising flatter areas surrounded by hilly areas. The laser beam diameter (~0.8 μ m) was sufficiently small to lie within these regions and so measurements were made in three areas on each sample: a flatter area, a hilly area, and an area combining both topological areas. Generally, the features observed in each spectrum were the same for all areas, but the hilly regions gave the strongest signal consistent with their greater film thickness (~1 μ m higher than the flatter areas). From depth dependent measurements of the Raman intensity in flatter areas using the confocal feature of the Raman microscope, the Ni(OH)₂ film thicknesses were estimated to be ~5 μ m. Some of the samples exhibited irregular cracks in the film (see, for example, Figure 1) that were probably

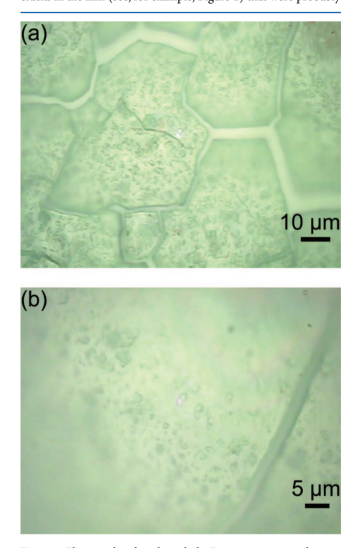


Figure 1. Photographs taken through the Raman microscope objective with objectives of (a) 50× and (b) 100× showing the cracked and bumpy nature (hence the fuzzy focus in regions of different height) of the Ni(OH)₂ film for sample α_3 .

introduced by stress during drying of the films. These cracks, which could reach down to the Ni substrate, were avoided during Raman measurements. Data were smoothed for presentation with a 25-point, second order Savitzky–Golay filter.

2.5. Fourier Transform-Infrared (FT-IR) Spectroscopy. The infrared measurements were performed at room temperature in a nitrogen-purged sample chamber using a Bruker Hyperion microscope with a 15× objective on a Bruker Tensor 27 FT-IR operated in the reflective mode and equipped with a mercury cadmium telluride (MCT) detector. The light diameter was several millimeters and thus included both topological areas described above. An untreated substrate was used to set the instrument zero. Spurious peaks arising from H_2O and CO_2 in the sample chamber were removed from the spectra with OPUS software (v.4.2, Bruker Optik GmbH).

3. RESULTS AND DISCUSSION

3.1. X-ray Diffraction Patterns. All measured XRD patterns show three intense peaks that arise from the metallic Ni substrate, thus indicating that the X-rays penetrate the entire film (Figure 2). The relative intensities of the Ni metal (220) peaks in

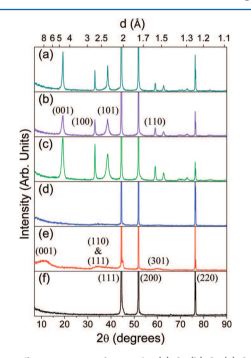


Figure 2. Diffraction patterns for samples (a) β_{11} (b) β_{22} (c) β_{33} (d) α_{22} (e) α_{33} and (f) Ni metal substrate with the Miller indices for the first diffraction peaks of each material identified. All patterns are plotted on the same intensity scale. A Cu K α X-ray source was used.

the β -Ni(OH)₂ samples are much weaker than for the α films. This indicates that the transmission at this angle is greater for the α than for the β films, implying that the β films are more crystalline than the α films and, from the relative intensities between samples, qualitatively shows that the sample thickness is $\beta_1 > \beta_2 > \beta_3$. The patterns from the β -Ni(OH)₂ films show all the expected peaks for this phase.⁴⁴ The (100) peaks are sharper than the (001) and (101) peaks, which indicates that there is greater order within the *ab*-plane than along the *c*-direction. This observation, with the layered structure of β -Ni(OH)₂, implies stacking faults of uniform Ni(OH)₂ sheets. From the full-width half-maxima of the (100) peaks, the β -films have similar crystallinity within the *ab*-plane, whereas the disorder in the *c*-direction is the greatest for sample β_{33} and the least for sample β_1 . The weak, broad diffraction peaks in the α -Ni(OH)₂ samples demonstrate that these are highly disordered films, which is expected from previous studies that report similar diffraction patterns for α -Ni(OH)₂.¹⁰ Only the Ni substrate was clearly observed in the XRD pattern for sample α_1 because the samples were thinner and had a high degree of structural disorder.

3.2. X-ray Photoelectron Spectra. The XPS measurements of samples β_1 , β_2 , and β_3 (see Figure 3a) agree well with reported

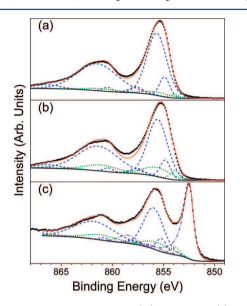


Figure 3. X-ray photoelectron spectra (\blacksquare) for samples (a) β_3 , (b) α_3 , and (c) Ni metal substrate curve-fitted with β -Ni(OH)₂ (---), NiO (•••), and Ni (-•-) components using the line shapes and method of Biesinger et al.⁵⁵ No literature references are presently available for the line shape of α -Ni(OH)₂. The solid red line shows the calculated line shape, and the solid black line shows the Shirley background.

 β -Ni(OH)₂ spectra.⁵⁵ Sample β_1 shows that some K (~2%) from the KOH aging solution remained on the surface. β_2 is pure, and β_3 contains a small amount of Ca (~0.7%), likely from impurities present in the Ni(NO₃)₂ salt used to deposit this film. Sample α_3

was prepared twice, and Ca was detected in only one of the films (~0.2%). Sample α_1 contains S (~2%), α_2 contains Cl (~2%), and sample α_3 contains N (~3%). This suggests that impurities, i.e., sulfate, chloride, and nitrate anions, are incorporated during electrochemical precipitation of α -Ni(OH)₂ and are eliminated from the structure during the aging from the α to the β form.

The Ni 2p region for the α samples shows Ni^{II} peaks similar to those observed in the β -Ni(OH)₂ spectra. However, subtle differences in the peak shapes and relative intensities between the α and β phases result in greater deviations between fitted lineshapes and the measured data (Figure 3b). At present, there are no reported XPS peak-fitting parameters for α -Ni(OH)₂.

These measurements demonstrate that XPS cannot easily differentiate the two phases. However, XPS can differentiate $Ni(OH)_2$ from metallic Ni and other nickel oxides, such as the air-formed nickel oxide on the Ni substrate (Figure 3c). It is also an effective way to identify and quantify elemental impurities on the surface.

3.3. Raman Spectra. As measured, several of the Raman spectra show a strong background (Figure 4i) that can be removed computationally (Figure 4ii). All samples show a sharp feature at 2330 cm⁻¹, which is an artifact of the laser excitation source. All samples also show a weak feature at ~1556 cm⁻¹, which corresponds to a two-magnon transition that arises from the room temperature antiferromagnetism of NiO.⁵⁶ The presence of this peak demonstrates that the air-formed oxide (NiO_x, $x \approx 1$) on the metal substrate, only 9–12 Å thick, ^{17,18} underlies all the deposited Ni(OH)₂ films. The Raman spectra of samples β_1 , β_2 , and β_3 are very similar, whereas the spectra of sample α_3 contain several peaks that are not observed for sample α_2 .

The Raman spectra for sample α_1 (not shown) have very low signal-to-noise and only contain a broad, weak band centered at ~3575 cm⁻¹ (α -Ni(OH)₂, O–H stretch) and medium peaks from sulfate ions (see section 3.8). Because the weak lattice modes could not be observed, this sample is not discussed in more detail.

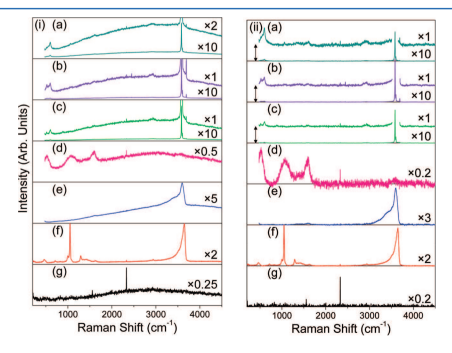


Figure 4. Raman spectra of samples (a) β_1 , (b) β_2 , (c) β_3 , (d) α_2 , (e) α_3 , and (f) the untreated Ni substrate. The data are shown on the same intensity scale (i) as collected and (ii) after background subtraction. The *y*-axis is linear, and select traces are rescaled by a factor, indicated on the plots. Traces iia, iib, and iic are offset as shown by double-headed arrows.

3.4. Infrared Spectra. All IR spectra contain sharp, weak features corresponding to the asymmetric bending, at 668 cm⁻¹, and asymmetric stretching, at 2341 cm⁻¹ and 2361 cm⁻¹, modes of CO₂ gas and some sharp, weak interference that arises from bending, at ~1600 cm⁻¹, and asymmetric stretching, at ~3750 cm⁻¹, modes of H₂O vapor in the sample chamber (Figure 5). All

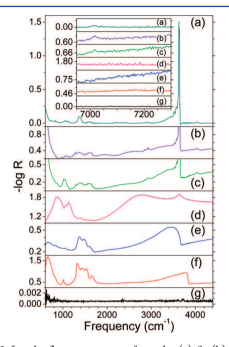


Figure 5. Infrared reflectance spectra of samples (a) β_1 , (b) β_2 , (c) β_3 , (d) α_1 , (e) α_2 , (f) α_3 , and (g) Ni substrate. The broad feature in spectra d at ~2600 cm⁻¹ is an artifact of the Fourier transform from low sample reflectance. Inset shows high frequency combination bands. Inset spectra are all plotted on the same $-\log R$ scale with each box 0.06 units in height and tick marks spaced 0.03 units apart.

spectra also contain a sharp, weak feature at 2170-2180 cm⁻¹ due to an adsorbed species, likely CO₂. The absolute absorbance of each sample cannot be determined because the exact film thicknesses are unknown, which affects the relative peak intensities. The reflectance of the substrates varies slightly. This causes the spectra to be offset by unknown, however, constant values, assuming that the reflectance is independent of the wavelength. A large, periodic background arising from the Fourier transform procedure was computationally removed from the IR spectra of sample β_1 . Therefore, peak shapes and absolute reflectance values are subject to some error. All other IR spectra are presented as measured. The results of the IR measurements complement the Raman spectra well, showing that samples β_1 , β_2 , and β_3 are very similar, whereas sample α_3 shows several peaks that are not observed in samples α_1 or α_2 , which correspond to nitrate bands. Sample α_1 has very high reflectance and a low signal-to-noise ratio, which indicates that it is very disordered and scatters the incident IR beam. Several peaks that are not present in samples α_2 and α_3 are observed in α_1 that originate from sulfate anions.

The IR spectra for β -Ni(OH)₂ are more complex than the Raman spectra because of transverse optical (TO)/longitudinal optical (LO) splitting of the vibrational modes. For asymmetric vibrational modes involving ionic species, TO modes do not create long-range Coulomb effects, whereas LO modes do. Additional restoring forces act upon the oscillating nuclei and cause LO transitions to occur at higher energies than their

corresponding TO transitions. Further, LO and TO transitions of different modes with the same symmetry may not cross over in energy.⁵⁷ The TO/LO splitting also causes peak intensities to depend on the crystal orientation relative to the polarization of the IR beam. However, this study examines polycrystalline samples using nonpolarized light, so this effect is not observed. The LO/TO effect is not observed for symmetric, e.g., Ramanactive, modes.

3.5. Vibrational Modes of β -Ni(OH)₂. *3.5.1. Raman-Active Vibrational Modes of* β -Ni(OH)₂. From the symmetry of β -Ni(OH)₂, the factor group theory predicts four Raman-active transitions, two with E_g symmetry and two with A_{1 g}.⁵⁷ It is well-established that three of these modes are at 310–315 cm⁻¹, 445–453 cm⁻¹, and 3581 cm⁻¹. The symmetry of these transitions has been disputed because the position of the fourth mode has not been clearly determined.⁴¹ However, it is worth noting that the Raman-active lattice modes of brucite [Mg(OH)₂], which is isostructural, are at similar positions: 280 cm⁻¹, 443 cm⁻¹, 725 cm⁻¹, and 3652 cm^{-1.58} A detailed computational analysis of brucite shows the atomic displacements corresponding to each mode.⁵⁷ Thus, the peak measured at ~880 cm⁻¹ (Figure 6a) is

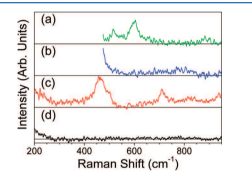


Figure 6. Raman spectra of samples (a) β_3 , (b) α_2 , and (c) α_3 showing lattice modes. (d) The Raman spectrum of water is shown for comparison. All spectra are plotted on the same intensity scale.

here ascribed to the final lattice mode of β -Ni(OH)₂ and the symmetries of the transitions can be assigned (Tables 2–5).

An additional band at $508-519 \text{ cm}^{-1}$ is sometimes reported and is visible in Figure 6a. The intensity of this feature varies between samples and is generally more intense for less crystalline samples.⁴¹ Although it has sometimes been ascribed to a lattice mode, the frequency is much lower than expected for the E_g mode, ⁵⁷ and its intensity can vary independently of the other lattice modes.³⁰ Inelastic neutron scattering measurements show that there is an acoustic vibrational mode at ~250-270 cm⁻¹, the harmonic overtone of which should be observable by Raman spectroscopy.⁴¹ Second order modes are typically very weak and increase in intensity for disordered samples; hence, the peak is assigned to this transition, consistent with the reported intensitycrystallinity correlation.

A Raman band is observed at 601 cm⁻¹ (Figure 6a) that has not been previously reported, presumably due to instrumental limitations. Since this peak is very weak and is not a predicted lattice mode, this peak is taken to be the harmonic overtone of the E_{σ} transition at 306–318 cm⁻¹.

A very weak feature at ~1600 cm⁻¹ is superimposed with a weak peak at ~1630 cm⁻¹ (Figure 7c). The latter is ascribed to the O–H bend of H₂O that has adsorbed on the material surface or has been trapped within the material, whereas the former is similar in shape to free water, however, in the wrong position.

Table 2. Raman Peaks for β -Ni(OH)₂ Prepared from Ni(NO₃)₂ Then Aged in KOH $[\beta_3]^a$

	peak equency (cm ⁻¹)	width	peak intensity	peak ass	ignment
	516	medium	weak	β -Ni(OH) ₂	2nd order acoustic mode
	601	medium	medium	β -Ni(OH) ₂	2 E _g
	$\sim 880^{b}$	broad	very weak	β -Ni(OH) ₂	E _g O–H bend
	1055 ^c	medium	weak	SO4 ²⁻	ν_1
	1556	very sharp	extremely weak	air-formed NiO _x	2 magnon
	~1600	broad	very weak	trapped H ₂ O	O–H bend
	~1635	broad	weak	free H ₂ O	O–H bend
	~2900	broad	very weak	surface hydrocarbon	C–H stretch
	3581	sharp	very strong	β -Ni(OH) ₂	A _{1g} O–H stretch
	3601	sharp	strong	disordered β - Ni(OH) ₂	O-H stretch
	3655 ^d	sharp	very weak	disordered β- Ni(OH) ₂	O-H stretch
	3688	sharp	weak	disordered β -Ni(OH) ₂	O–H stretch, surface
-					

^{*a*}Samples aged in KOH after preparation from NiSO₄ [β_1] and NiCl₂ [β_2] gave the same results except as noted. ^{*b*}Not observed in samples prepared from NiSO₄ [β_1]. ^{*c*}Only observed in samples prepared from NiSO₄ [β_1]. ^{*d*}Only observed in samples prepared from Ni(NO₃)₂ [β_3].

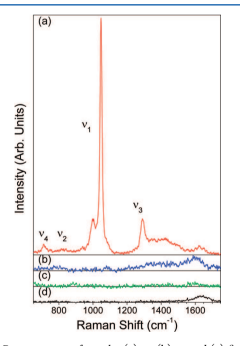


Figure 7. Raman spectra of samples (a) α_3 , (b) α_2 , and (c) β_3 showing O–H internal modes. The nitrate ion vibrational modes, labeled $\nu_1 - \nu_4$ are shown for sample α_2 . These features are absent in samples α_1 and β_2 . (d) The Raman spectrum of water is shown for comparison. All spectra are plotted on the same intensity scale.

Thus, this is taken to be the O–H bend for a small degree of hydration in the structure (formally termed a libration mode for lattice species⁵⁷).

A sharp band at 3601 cm⁻¹ is present in the Raman spectra for samples β_1 and β_2 . This peak has previously been variously assigned to adsorbed H₂O, crystal defects, impurities, surface OH groups, and the O–H stretch for a new crystal phase, β_{bc} -Ni(OH)₂.^{29,31,41} The intensity of this peak has been reported to increase with decreasing crystallite size, as measured by XRD peak width.²⁹ However, only the diffraction peaks involving the crystallographic *c*-axis were broadened. This indicates stacking faults rather than small crystallites, and thus, the Scherrer equation cannot be properly applied. Re-examination of the results reveals that the Raman peak intensity at 3601 cm⁻¹ increases as disorder along the *c*-axis increases. The peak intensity decreases when samples are heated to 100–150 °C.²⁹

We now propose that the $\beta_{\rm bc}$ phase is more accurately described as β -Ni(OH)₂ that has stacking fault disorder. In the O-H stretching modes, the H atoms approach adjacent sheets more closely than in other vibrational modes, and hence, the Raman peak at 3581 cm⁻¹ is shifted to 3601 cm⁻¹. There must be a finite number of relative sheet positions and orientations that occupy local free energy minima. If the sheets do not rotate, then two structures are predicted, namely (a) Ni atoms in register between adjacent sheets, which is the thermodynamic minimum, and (b) a staggered configuration in which Ni atoms are in register between alternating sheets (if the sheets do rotate, then the rotation would be by 60° , and hence, two more new structures are permitted). Though the true geometry is unknown, this kinetically stable structure has slightly different O-H stretching modes and restructures to the stable form when heated to 100-150 °C. This disordered structure has never been reported in excess of the ordered β -Ni(OH)₂ component, and thus, it can only coexist as part of a parent β phase material. Therefore, the evidence does not support the existence of an additional crystallographic phase but rather a disordered form of β -Ni(OH)₂.

The number of exchanged electrons per Ni atom in β -Ni(OH)₂ electrodes has been found to approach unity for materials with a strong Raman band at 3605 cm⁻¹ (corresponding to our peak at 3601 cm⁻¹), whereas it is close to 0.8 for samples that do not show this peak. This was previously attributed to the increased surface area of smaller crystallites.²⁹ However, our reinterpretation of this peak leads to the conclusion that materials with *c*-axis disorder are better candidates than more ordered structures. The photocatalysts prepared by Yu et al. also have *c*-axis disorder that can be observed from their XRD patterns,¹⁵ which may indicate that photocatalytic activity is also linked to stacking faults in the crystal structure.

A weak but sharp peak is observed at 3655 cm⁻¹ that is assigned to the IR-active disordered O–H stretch. Because this is an out-of-phase peak, it is quite weak in the Raman spectrum. However, the disorder or the change in symmetry makes it only weakly visible. It is also possible that this is a localized mode around Ca point defects from the Ni(NO₃)₂ starting material (cf., A_{1g} for bulk Ca(OH)₂ is observed at 3620 cm⁻¹⁵⁸). Ca was not detected in the more-ordered samples β_1 and β_2 , and the peak is not present in their Raman spectra, which makes either explanation equally plausible.

One additional Raman peak is observed at 3688 cm⁻¹. The intensity is sometimes reported to be linked to that of the peak at 3601 cm^{-1} . However, the peaks have also been observed to vary independently of each other.^{29,30} Bernard et al. observed that dry samples, which initially show this peak, that are rinsed with water and reanalyzed while still wet, lose the peak completely.³⁰ This indicates that it is a surface-related phenomenon, although the exact origin is unclear. Bernard et al. proposed that the O–H stretch for surface hydroxide groups is at higher energy than for those opposed by adjacent layers. Upon hydration, the transition becomes similar to either the bulk O–H stretching mode, A_{1e} , or

that of free H_2O .³⁰ The peak observations and assignments are summarized in Table 2.

3.5.2. Infrared-Active Vibrational Modes of β -Ni(OH)₂. Factor group theory predicts four IR-active vibrational modes for β -Ni(OH)₂: Two A_{2u} and two E_u.⁵⁷ There are twice as many IR peaks as there are Raman peaks for β -Ni(OH)₂ because of transverse optical (TO)/longitudinal optical (LO) splitting (see section 3.4). The magnitude of this splitting effect has been calculated for brucite and can be used to assist peak identification for β -Ni(OH)₂.⁵⁷

Only four IR lattice modes have been previously identified: 332-354 cm⁻¹, 440-475 cm⁻¹, 510-553 cm⁻¹, and 3630- $3650 \text{ cm}^{-1.41}$ The LO/TO splitting for the A_{2u} mode at ~3635 cm⁻¹ is expected to be quite small (cf., 14 cm⁻¹ for Mg(OH)₂⁵⁷). The observed peaks for samples β_2 and β_3 exhibit flat tops because of the low reflectance at this higher sample thickness. The peak for the thinner sample β_1 is better defined and shows peaks at 3629 cm⁻¹ and 3643 cm⁻¹, which are assigned to the A₂₁ (TO) and $A_{2\mu}$ (LO) modes, respectively. An IR study of β -Ni(OD)₂ showed that the two apparent peaks at 350 cm⁻¹ and 450 cm⁻¹ are actually each composed of two overlapping peaks: One each that red-shifts on deuteration and one each that does not observably shift. The peak at 550 cm⁻¹ is also observed to red-shift on deuteration.⁷ The two remaining A_{2u} modes are not predicted to significantly shift on deuteration, whereas all four of the Eu modes are;⁵⁷ hence, these peaks can be assigned accordingly. This is consistent with a study that observed the shift in position of three lattice modes for $Mg_{1-x}Ni_x(OH)_2$, where $0 \le x \le 1.^{20}$ The remaining E_u mode is predicted at a frequency greater than 550 cm⁻¹ and has a very large TO/LO split (cf., 370 cm^{-1} for brucite); thus, this mode is assigned to the broad band at $\sim 1030 \text{ cm}^{-1}$ (Figure 9). Because of its low intensity, this peak is previously unreported, although close examination reveals that it may be visible as a weak shoulder in published results.²⁹ Thus, the eight IR-active lattice modes have now been identified and assigned (Table 5).

A broad peak at ~1630 cm⁻¹ (Figure 9) arises from the O–H bending mode of H_2O that is either adsorbed on the material surface or trapped within the structure. The peaks at ~3380 cm⁻¹ and ~3450 cm⁻¹ are from the O–H stretch of the H_2O (Figure 10).

A sharp but weak peak measured at 3581 cm⁻¹ corresponds exactly to the Raman-active A_{1g} transition (Figure 10). This is formally forbidden in IR spectroscopy; however, structural disorder allows a very weak peak to be seen in the IR spectra. Conversely, a weak A_{2u} mode should also be allowed in the Raman spectrum. However, this mode overlaps with other features of the sample and hence cannot be observed in the Raman spectrum. A shoulder at 3601 cm⁻¹ corresponds to the inphase O–H stretch of disordered β -Ni(OH)₂ discussed in the Raman section. The corresponding out-of-phase O–H stretch (LO) is a strong, sharp peak at 3652 cm⁻¹. The corresponding TO mode is visible in the spectra for sample β_1 as a shoulder at 3634 cm⁻¹. Note that disorder shifts both the TO and LO modes to higher frequency by 5–9 cm⁻¹, which reaffirms that these two peaks are connected.

The peak at 2658 cm⁻¹ is sharp, very weak, and close to the positions reported for the A_{2u} mode of β -Ni(OD)₂ (cf., 2705 cm⁻¹,⁷ 2680 cm⁻¹,²⁵ and 2690 cm⁻¹⁴²). The difference in position could arise because this study observes localized modes from the naturally present D (0.0115%), whereas the reference materials were highly enriched.

Several previously unreported weak features are observed in the IR spectra for β -Ni(OH)₂ that can be ascribed to combination bands. Since this structure has an inversion center, the overall symmetry must be odd (ungerade) for a transition to be IR-active. Thus, all overtone transitions are even and therefore Raman-active, whereas an IR mode or an acoustic mode combined with a Raman mode is odd and therefore IR-active. The peak assignments (Tables 3 and 4) were created by

Table 3. Infrared Peaks for β -Ni(OH)₂ Prepared from NiSO₄ Then Aged in KOH [β_1]

peak frequency (cm ⁻¹)	width	peak intensity	peak	assignment
<600 ^a	medium	strong	β -Ni(OH) ₂	A _{1g} + acoustic mode combination
~635 sh	medium	weak	β -Ni(OH) ₂	$\begin{array}{c} E_{g}+A_{2u} (TO)/E_{g}+E_{u} \\ (TO) \text{ combination} \end{array}$
~1045	medium	medium	β -Ni(OH) ₂	E_u (LO) O–H bend
1285	sharp	weak	surface hydrocarbon	C–H bend
1385	medium	medium	β -Ni(OH) ₂	$E_{g} + A_{2u}$ (TO) combination
~1560	medium	weak	β -Ni(OH) ₂	$E_u (LO) + 2nd order$ acoustic mode
~1640	medium	weak	free H ₂ O	O–H bend
2655	sharp	very weak	β -Ni(OH) (OD)	A _{2u} (TO/LO) O–D stretch
2855	medium	weak	surface hydrocarbon	C–H stretch
2880 sh	medium	weak	surface hydrocarbon	C–H stretch
2928	medium	weak	surface hydrocarbon	C–H stretch
2960 sh	medium	weak	surface hydrocarbon	C–H stretch
3380	medium	weak	free H ₂ O	O–H stretch
3450	medium	weak	free H ₂ O	O–H stretch
3520	sharp	weak	β -Ni(OH) ₂	A _{1g} – acoustic mode subtraction
3566	sharp	weak	β -Ni(OH) ₂	A _{1g} – acoustic mode subtraction
3580	sharp	medium	β -Ni(OH) ₂	A _{1 g} O–H stretch
3601 sh	sharp	medium	disordered β -Ni(OH) ₂	O—H stretch (in- phase)
3629	sharp	very strong	β -Ni(OH) ₂	A _{2u} (TO) O–H stretch
3634 sh	sharp	strong	disordered β- Ni(OH) ₂	O–H stretch (TO, out-of-phase)
3643	sharp	very strong	β -Ni(OH) ₂	A _{2u} (LO) О–Н stretch
3652	sharp	strong	disordered β - Ni(OH) ₂	O–H stretch (LO, out-of-phase)
4040	medium	medium	β -Ni(OH) ₂	$\begin{array}{c} A_{1g} + A_{2u} (LO) / A_{1g} + \\ E_{u} (LO) \\ \text{combination} \end{array}$
4270	broad	very weak	β -Ni(OH) ₂	3rd order combination
7020	sharp	very weak	β -Ni(OH) ₂	$A_{1g} + A_{2u}$ (TO/LO) combination

 ${}^{a}\mbox{Peak}$ maximum is below the instrumental cutoff, so the exact position is unknown.

considering the IR and Raman lattice modes (Table 5) and the acoustic transitions observed by INS.⁴¹ To conclusively assign these peaks, single crystal measurements using plane-polarized light are necessary, especially the modes for which multiple combinations are possible. One particularly interesting combination is the sharp peak at 3520 cm⁻¹ assigned to the

Table 4. Infrared	Peaks for β -Ni(OH)	² Prepared from
$Ni(NO_3)_2$ Then A	Aged in KOH $[\beta_3]^a$	

peak frequency cm ⁻¹	width	peak intensity	peak ass	ignment
<600 ^b	medium	strong	β -Ni(OH) ₂	A _{1g} + acoustic mode combination
665 sh	medium	weak	β -Ni(OH) ₂	$\begin{array}{c} E_g + A_{2u} (TO) / E_g + \\ E_u (TO) \\ combination \end{array}$
1030	medium	medium	β -Ni(OH) ₂	E_u (LO) O–H bend
1285	sharp	weak	surface hydrocarbon	C–H bend
~1400	broad	medium	β -Ni(OH) ₂	$E_{g} + A_{2u}$ (TO) combination
~1550	medium	medium	β -Ni(OH) ₂	$E_u(LO) + 2nd order$ acoustic mode
~1630	broad	medium	free H ₂ O	O–H bend
2658	sharp	very weak	β -Ni(OH)(OD)	A _{2u} (TO/LO) O–D stretch
2855 sh	medium	weak	surface hydrocarbon	C-H stretch
2880	medium	weak	surface hydrocarbon	C-H stretch
2930	medium	weak	surface hydrocarbon	C-H stretch
2960 sh	medium	weak	surface hydrocarbon	C-H stretch
3140	medium	very weak	β -Ni(OH) ₂	O–H bend combination
3380	medium	weak	free H ₂ O	O–H stretch
3450	medium	weak	free H ₂ O	O-H stretch
3520	sharp	weak	β -Ni(OH) ₂	A _{1g} – acoustic mode subtraction
3566	sharp	very weak	β -Ni(OH) ₂	A_{1g} – acoustic mode subtraction
3581	sharp	weak	β -Ni(OH) ₂	A _{1g} O–H stretch
3601 sh	sharp	very weak	disordered β- Ni(OH) ₂	O–H stretch (in- phase)
~3635	sharp	extremely strong	β -Ni(OH) ₂	A _{2u} (TO/LO) O–H stretch
3652	sharp	strong	disordered β- Ni(OH) ₂	O-H stretch (LO, out-of-phase)
4035	medium	weak	β -Ni(OH) ₂	$\begin{array}{l} A_{1g} + A_{2u} \; (LO) / A_{1g} \\ + \; E_{u} \; (LO) \\ \text{combination} \end{array}$
~4275	broad	very weak	β -Ni(OH) ₂	3rd order combination
7020	sharp	very weak	β -Ni(OH) ₂	$\begin{array}{c} A_{1g} + A_{2u} \left(TO/LO \right) \\ combination \end{array}$

"The sample prepared from NiCl₂ then aged in KOH $[\beta_2]$ gave the same results. ^{*b*}Peak maximum is below the instrumental cutoff, so the exact position is unknown.

combination of the A_{1g} mode and a subtractive mode, i.e., a hot band, for the acoustic transition at ~60 cm⁻¹ (Figure 10).⁴¹ Hot bands are not typically observed in IR spectra at room temperature. However, the acoustic transition is sufficiently low in energy that a Boltzmann distribution predicts a population ratio of 57 to 43 for the ground state and first excited state, respectively. Another combination of the A_{1g} and a subtractive mode is observed at 3566 cm⁻¹.

3.6. Vibrational Modes of α -Ni(OH)₂. *3.6.1. Raman*-Active Vibrational Modes of α -Ni(OH)₂. The Raman spectra of α -Ni(OH)₂ show lattice modes at 460 cm⁻¹ and 495 cm⁻¹ (Figure 6). Presumably due to instrumental limitations, only the former peak has been previously reported.^{32,34,36} Weak features, assigned in this work to second order transitions, are observed at \sim 790 cm⁻¹ and \sim 1075 cm⁻¹. Some authors report additional Raman-active lattice modes.^{32,39} However, these are not observed in the present work and can all be ascribed to β -Ni(OH)₂ impurities. Structural disorder causes the internal O–H bending mode of lattice OH to split and broaden into a weak feature at ~1400 cm⁻¹ (Figure 7). Since the Raman intensity is quite weak, it is necessary to use IR, discussed below, to observe fine details for these peaks. Further, this broad feature underlies several nitrate bands in sample α_3 and is therefore unclear. A second internal O–H bending mode at 1590–1620 cm⁻¹ is similar in width and only slightly shifted from the position of free H₂O. Thus, this can be assigned to the O–H bending mode of intercalated H₂O. A third internal O–H bending mode is observed at 1630 cm⁻¹, which arises from adsorbed H₂O on the surface or H₂O trapped within the structure.

There are several O-H stretching modes from 3240 to 3515 cm⁻¹ that arise from free H₂O (Figure 8). The internal O-H stretching modes from lattice OH and intersheet H₂O are visible from 3590 to 3650 cm^{-1} . This region is quite different between samples α_2 and α_3 and such differences have previously been suggested to arise from the presence of intermediate phases.³⁶ However, since the lattice modes do not vary as significantly between samples, this supports that this phase tends to adopt disorder by the incorporation of foreign ions, variable hydration, and randomly oriented layers.¹¹ This is confirmed by our measurement of two preparations of sample α_2 where no variation in lattice mode frequencies was found, but the frequencies of the internal modes of the nitrate ions (see section 3.7) and the O-H stretching frequency are different. The shifts in frequency are caused by variations in the local environment and/or stress (see section 2.4). These peak observations and assignments are summarized in Tables 6 and 7.

3.6.2. Infrared-Active Vibrational Modes of α -Ni(OH)₂. There are three IR-active α -Ni(OH)₂ modes at 380–400 cm⁻¹, 460–480 cm⁻¹, and 625–670 cm⁻¹ (Figure 9). Only the last of these is observed in this work due to instrumental limitations. Weakly visible combination bands are observed at 835 cm⁻¹ and 1055 cm⁻¹. The latter is not visible in the spectra of sample α_2 because it is obscured by nitrate bands.

Two peaks from the O–H bending modes of lattice OH are observed at 1380–1390 cm⁻¹ and 1480–1490 cm⁻¹ (Figure 9). These peaks shift slightly between samples because of structural disorder and the mechanical stress induced when the materials dried. The O–H bending mode from intercalated H₂O is measured at 1600 cm⁻¹, which matches the Raman spectra. Note that the absorbance of the first three O–H bending modes is too great to be assigned to free H₂O that is trapped within the material, whereas a weaker peak is observed at 1630 cm⁻¹ that is consistent with this assignment.

As in the Raman spectra, several O–H stretching modes are observed in the IR spectra of the α phase samples (Figure 10). Broad features from free H₂O are measured at 3220–3515 cm⁻¹, and peaks corresponding to lattice OH and intersheet H₂O are observed at 3570–3640 cm⁻¹. The variability from structural disorder is the same as in the Raman spectra.

A combination band of the O–H stretch and a lattice mode is measured at 4260–4280 cm⁻¹. A combination of O–H stretching modes is measured at 7140 cm⁻¹ (Figure 5 inset). Additional features are difficult to distinguish by FT-IR because of periodic interference fringes arising from the Fourier transformation. These peak observations and assignments are summarized in Tables 8–10. Our results are compared with literature values in Table 11.

3.7. Nitrate Bands. From the XPS elemental analysis, nitrate bands are only expected in the vibrational spectra of sample α_2 . Free nitrate has D_{3h} symmetry and four vibrational modes: ν_1 at

Table 5. Comparison of Literature Values for the Vibrational Modes in β -Ni(OH)₂ and Equivalent Modes in Mg(OH)₂ (Brucite)

		Mode Frequency (cm ⁻¹)	
our peak assignment	this study	literature	$Mg(OH)_2$
Raman			
E _g lattice mode		315, ⁵ 319, ²² 310, ²⁹ 310, ³⁰ 312–313, ³² 318, ³⁵ 306, ³⁶ 315, ³⁸ 309, ³⁹ 313 ⁴¹	280 ⁵⁸
A _{1g} lattice mode		450, ⁵ 450, ¹⁹ 458, ²² 450, ²⁹ 445, ³⁰ 448–453, ³² 449, ³⁵ 445, ³⁶ 450, ³⁸ 446, ³⁹ 447 ⁴¹	443 ⁵⁸
2nd order acoustic mode	516	510, ¹⁹ 519, ²² 515, ²⁹ 510, ³⁰ 519, ³² 510, ³⁶ 517, ³⁸ 510, ³⁹ 508 ⁴¹	
2Eg 2nd order lattice mode	601		
E _g O–H bend	~880	~850, ³⁵ 880 ³⁸	725 ⁵⁸
trapped H ₂ O, O–H bend	~1600		
free H ₂ O, O–H bend	~1635		
A _{1g} O–H stretch	3581	3581, ⁵ 3599, ²² 3580, ²⁸ 3585, ²⁹ 3570, ³⁰ 3565, ³¹ 3580–3583, ³² 3580, ³⁵ 3579, ³⁶ 3580, ³⁸ 3579, ³⁹ 3581 ⁴¹	3652 ⁵⁸
disordered, O–H stretch	3601	3601, ⁵ 3605, ²⁹ 3580, ³⁰ 3600, ³¹ 3600, ³² 3600, ³⁸ 3599, ³⁹ 3601 ⁴¹	
surface, O–H stretch	3688	3680, ³⁰ 3687, ³¹ 3687 ³²	
Infrared			
A_{2u} (TO)/ E_u (TO) lattice modes		350, ⁶ 350, ⁷ 348, ²⁰ 345, ²¹ 340, ²² 350, ²⁴ 354, ²⁵ 350, ³⁵ 332 ⁴¹	368 ⁶³
A_{2u} (LO)/ E_u (LO) lattice modes		~450, ¹ 450, ⁶ 450, ⁷ 451, ²⁰ 460, ²¹ 458, ²² 460, ²⁴ 475, ²⁵ 460, ²⁹ 452, ³⁵ 440 ⁴¹	562 ⁶³
E _u (TO) O–H bend		517, ¹ 530, ⁶ 545, ⁷ 549, ²⁰ 520, ²¹ 525, ²² 510, ²⁴ 530, ²⁵ 525, ²⁹ 553, ³⁵ 530 ⁴¹	462 ⁶³
E _u (LO) O–H bend	1030	1030 sh ²⁹	
free H ₂ O, O–H bend	~1630	1620, ⁶ 1630 ²⁹	
A_{2u} (TO) O–H stretch	3629	3640, ¹ 3650, ⁶ 3650, ⁷ 3650, ¹⁴ 3635, ²⁰ 3650, ²¹ 3650, ²⁴ 3650, ²⁵ 3639, ²⁸ 3630, ³⁵ 3636, ⁴¹ 3640 ⁴²	3698 ⁶³
disordered, (TO) O–H stretch	3634		
A _{2u} (LO) O–H stretch	3643		
disordered, (LO) O–H stretch	3652		
$A_{1g} + A_{2u} (LO) / A_{1g} + E_u (LO)$	4035	4028 ²⁸	
3rd order combination	~4275	4274 ²⁸	
$A_{1g} + A_{2u} (TO/LO)$	7020	7016 ²⁸	

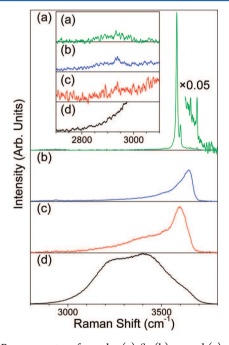


Figure 8. Raman spectra of samples (a) $\beta_{3^{j}}$ (b) $\alpha_{2^{j}}$ and (c) α_{3} showing O–H internal modes. (d) The Raman spectrum of water is shown for comparison. All spectra, except where noted, are plotted on the same intensity scale. Inset shows low frequency region of the figure enlarged by a factor of 15.

1050 cm⁻¹ (A'₁, IR-inactive), ν_2 at 831 cm⁻¹ (A"₂, Ramaninactive), ν_3 at 1390 cm⁻¹ (E'), and ν_4 at 720 cm⁻¹ (E').⁵⁹ In solution, the ν_1 band becomes weakly IR-active and the ν_3 band splits into a doublet at 1356 and 1410 cm⁻¹, both of which are weakly Raman-active. This split results from rotation and is

Table 6. Raman Peaks for α -Ni(OH)₂ Prepared from NiCl₂ [α_2]

peak frequency (cm ⁻¹)	width	peak intensity	peak ass	signment
~790	medium	weak	α -Ni(OH) ₂	2nd order lattice mode
~1075	medium	very weak	α -Ni(OH) ₂	2nd order lattice mode
~1400	broad	weak	α -Ni(OH) ₂	O–H bend, lattice OH
1555	very sharp	extremely weak	NiO	2 magnon
~1590	broad	weak	α -Ni(OH) ₂	O–H bend, layer H ₂ O
1630 sh	medium	very weak	free H ₂ O	O–H bend
~2900	broad	very weak	α -Ni(OH) ₂ /surface hydrocarbon	2nd order O–H bend/C–H stretch
2935	sharp	very weak	surface hydrocarbon	C-H stretch
~3240	very broad	weak	free H ₂ O	O-H stretch
~3380	very broad	medium	free H ₂ O	O-H stretch
~3515	broad	medium	free H ₂ O	O-H stretch
3590	medium	strong	α -Ni(OH) ₂	O–H stretch
3620	medium	strong	α -Ni(OH) ₂	O-H stretch
3650 sh	medium	medium	α -Ni(OH) ₂	O–H stretch

therefore only expected if the chemical environment allows free rotation. Further, these bands can shift and split into additional modes when nitrate interacts with cations to form ion pairs.⁶⁰

The Raman spectra of sample α_2 show all four nitrate modes (Figure 7) with the most intense peak at 1047 cm⁻¹ (ν_1), and the IR spectra also show all four free nitrate peaks (Figure 9) with the strongest peak at 1340 cm⁻¹ (ν_2). The hydrated intercalation

Table 7. Raman Peaks for α -Ni(OH)₂ Prepared from Ni(NO₃)₂ [α_3]

peak frequency (cm ⁻¹)	width	peak intensity	peak as	ssignment
460	medium	weak	α -Ni(OH) ₂	lattice mode
495 sh	medium	weak	α -Ni(OH) ₂	lattice mode
712/719 ^a	medium	weak	NO ₃ ⁻	$ u_4 $
816/840 ^a	medium	weak	NO ₃ ⁻	ν_2
~935	medium	very weak	α -Ni(OH) ₂	2nd order lattice mode
997/1002 ^a	sharp	medium	NO ₃ ⁻	ν_1
$1047/1048^{a}$	sharp	strong	free NO ₃ ⁻	ν_1
1075 sh	medium	weak	α -Ni(OH) ₂	2nd order lattice mode
1291/1293 ^a	medium	medium	NO ₃ ⁻	ν_3
~1350	medium	very weak	free NO ₃ ⁻	ν_3
~1420	broad	weak	NO ₃ ⁻	$2\nu_4$
~1495	broad	weak	NO ₃ ⁻	$\nu_2 + \nu_4$
1554	very sharp	extremely weak	air-formed NiO _x	2 magnon
~1620	medium	weak	α -Ni(OH) ₂	O–H bend, layer H ₂ O
~1630	broad	weak	Free H ₂ O	O–H bend
~1695	medium	very weak	NO ₃ ⁻	$2\nu_2$
~2900	broad	very weak	α-Ni(OH) ₂ / surface hydrocarbon	2nd order O–H bend/C–H stretch
~2935	medium	very weak	surface hydrocarbon	C–H stretch
~3240	broad	very weak	free H ₂ O	O–H stretch
3515 sh	broad	medium	free H ₂ O	O–H stretch
3647/3642 ^a	medium	strong	α -Ni(OH) ₂	O–H stretch

 a Two samples were used in this study (see section 2.1), and for some peaks, different values were obtained, as reported in the table.

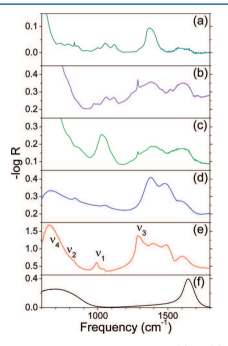


Figure 9. Infrared reflectance spectra of samples (a) β_1 , (b) β_2 , (c) β_3 , (d) α_2 , (e) α_3 , and (f) a thin film of water on a Ni substrate.

region is considered to be amorphous and can vary considerably in size, based on the variability of the *c*-dimension of the unit cell.¹⁰ Since these four nitrate modes are very similar to solution

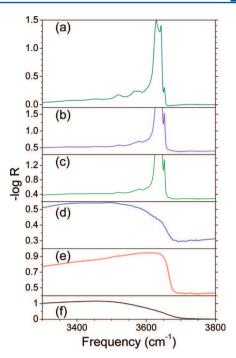


Figure 10. Infrared reflectance spectra of samples (a) β_1 , (b) β_2 , (c) β_3 , (d) α_2 , (e) α_3 , and (f) a thin film of water on a Ni substrate.

Table 8.	Infrared P	eaks for $\pmb{\alpha}$	-Ni(OH) ₂ I	Prepared	from
NiSO ₄ [α_1] ^{<i>a</i>}			-	

peak frequency (cm ⁻¹)	width	peak intensity	peal	x assignment
685 sh ^b	sharp	very weak	SO4 ²⁻	ν_2
830 sh	sharp	weak	α -Ni(OH) ₂	lattice mode combination
885	medium	medium	$\frac{SO_4^{2-}/\alpha}{Ni(OH)_2}$	$ u_4 $ + lattice mode combination
980 sh	medium	weak	free SO4 ^{2–}	$ u_1 $
1065	medium	medium	SO4 ²⁻	$ u_1 $
~1130	medium	medium	free SO4 ^{2–}	ν_3
1180 sh	medium	medium	SO4 ²⁻	ν_3
1410	medium	weak	α -Ni(OH) ₂	O–H bend, lattice OH
~1620	medium	weak	free H ₂ O	O–H bend
3650	broad	medium	α -Ni(OH) ₂	O–H stretch

^aTwo samples were used in this study (see section 2.1). ^bPeak maximum is near the instrumental cutoff, so the exact position may be lower.

nitrate, it is most likely that the nitrate ions are located in the interlayer space.

There are additional nitrate peaks close to the first three nitrate modes in the IR spectra, which implies that nitrate exists in two distinct chemical environments. It is noteworthy that the second ν_1 IR peak intensity is greater than the corresponding, hydrated nitrate peak, whereas in the Raman spectra, the additional mode is much less intense than the hydrated nitrate peak. This indicates that there are two sets of selection rules and confirms the presence of two types of nitrate, rather than peak splitting from ion-pairing effects. The change in selection rules also demonstrates that the second position in the crystal reduces the symmetry below D_{3h} . It is unclear whether the second type of nitrate has a different orientation within the interlayer space or whether it substitutes lattice hydroxide.

Table 9. Infrared Peaks for α -Ni(OH)₂ Prepared from NiCl₂ [α_2]

peak frequency (cm ⁻¹)	width	peak intensity	peak a	assignment
670 ^a	broad	strong	α -Ni(OH) ₂	lattice mode
835	sharp	medium	α -Ni(OH) ₂	lattice mode combination
1055	medium	medium	α -Ni(OH) ₂	lattice mode combination
1380	medium	strong	α -Ni(OH) ₂	O–H bend, lattice OH
1480	medium	strong	α -Ni(OH) ₂	O–H bend, lattice OH
1600 sh	medium	strong	α -Ni(OH) ₂	O–H bend, layer H ₂ O
1630 sh	medium	medium	free H ₂ O	O–H bend
2860	sharp	very weak	surface hydrocarbon	C–H stretch
2930	sharp	very weak	surface hydrocarbon	C–H stretch
3220 sh	broad	medium	free H ₂ O	O–H stretch
3380	broad	very strong	free H ₂ O	O–H stretch
3500	medium	strong	free H ₂ O	O-H stretch
3570	medium	very strong	α -Ni(OH) ₂	O-H stretch
3640 sh	medium	strong	α -Ni(OH) ₂	O-H stretch
4260	medium	very weak	α -Ni(OH) ₂	O–H stretch + lattice mode

^{*a*}Peak maximum is near the instrumental cutoff, so the exact position may be lower.

Previous authors have assigned the IR peaks differently.²⁷ However, the relative intensities of the four modes in the Raman and IR spectra confirms that the nitrate peaks are correctly identified in the present work.

From this work, the incorporation of nitrate anions into at least two distinct chemical environments in α -Ni(OH)₂ can been identified by examining the Raman and IR spectra. Thus, applied research on this material will now have a method to rapidly determine any links between changes in performance or stability and the inclusion of nitrate anions in these environments.

3.8. Sulfate Bands. From sample preparation and XPS elemental analysis, sulfate bands are expected only in the vibrational spectra of sample α_1 . The IR spectra show weak features at 980 cm⁻¹ and 1120 cm⁻¹ that are similar to the positions of the ν_1 and ν_3 modes of free sulfate (cf., 981 cm⁻¹ and 1104 cm⁻¹, respectively⁵⁹). By analogy to the nitrate bands discussed above, this may indicate that sulfate is in the hydrated interlayer space. A second pair of bands is more strongly observed in the Raman and IR spectra at 1060 cm⁻¹ and 1170 cm⁻¹, which suggests that sulfate ions are incorporated in some chemical environment that shifts the peaks to higher frequency.

Additional bands, corresponding to ν_2 and ν_4 , respectively, are measured by Raman at 520 cm⁻¹ and both IR and Raman at 685 cm⁻¹. These are slightly higher than expected for free sulfate (cf., 451 cm⁻¹ and 613 cm⁻¹, respectively⁵⁹); hence, they correspond to the peaks at 1060 cm⁻¹ and 1170 cm⁻¹. The Raman spectra also show a combination of ν_1 and ν_2 at 1590 cm⁻¹; however, this is not observed by IR. The high degree of structural disorder hinders further investigation, and it cannot presently be concluded whether some anions are incorporated into the α -Ni(OH)₂ structure.

Similar to the findings for nitrate anions, our work establishes a method to measure sulfate anions in α -Ni(OH)₂ using Raman or

Table 10. Infrared Peaks for α -Ni(OH)₂ Prepared from Ni(NO₃)₂ [α_3]

peak frequency (cm ⁻¹)	width	peak intensity	peak	assignment
650 ^a	broad	very strong	α -Ni(OH) ₂	lattice mode
720 sh	medium	very weak	NO ₃ ⁻	$ u_4$
805 sh	medium	weak	NO ₃ ⁻	ν_2
835	medium	medium	α -Ni(OH) ₂	lattice mode combination
992	medium	medium	NO ₃ ⁻	$ u_1 $
1042	medium	medium	free NO ₃ ⁻	$ u_1 $
1280	medium	strong	NO ₃ ⁻	ν_3
1310 sh	medium	strong	NO ₃ ⁻	ν_3
~1340	medium	strong	free NO ₃ ⁻	ν_3
~1390	medium	strong	α -Ni(OH) ₂	O–H bend, lattice OH
~1490	medium	strong	α -Ni(OH) ₂	O–H bend, lattice OH
~1600	medium	strong	α -Ni(OH) ₂	O–H bend, layer H ₂ O
1630 sh	medium	medium	free H ₂ O	O–H bend
~1755	medium	weak	α -Ni(OH) ₂ + NO ₃ ⁻	lattice mode + ν_3
~1970	medium	weak	α -Ni(OH) ₂ + NO ₃ ⁻	lattice mode + ν_3
~2485	medium	very weak	NO ₃ ⁻	$2\nu_3$
~2775	medium	weak	NO ₃ ⁻	$2\nu_3$
~3240	broad	medium	free H ₂ O	O–H stretch
3390 sh	broad	very strong	free H ₂ O	O–H stretch
3515 sh	broad	strong	free H ₂ O	O–H stretch
3600-3630	medium	very strong	α -Ni(OH) ₂	O–H stretch
4280	medium	weak	α -Ni(OH) ₂	O–H stretch + lattice mode
7140	medium	very weak	α -Ni(OH) ₂	O–H stretch combination

^{*a*}Peak maximum is near the instrumental cutoff, so the exact position may be lower.

IR spectroscopy. This will allow future studies to correlate the presence of sulfate anions to changes in behavior of an electrode or catalyst.

3.9. Fluorescence. The strong background signal observed in the Raman spectra of both β films and sample α_1 (Figure 4i) indicates that the samples are fluorescent. Fluorescence is also observed from the air-formed NiO_x on the substrate, but it is relatively weak. The fluorescence arises from the electronic structure of the Ni^{II} cations, which have 3d⁸ valence configurations and octahedral coordination of OH⁻ anions. Ideally, in the ground state (³A_{2g}) the three highest occupied orbitals have t_{2g} symmetry, and there are unpaired, spin-aligned valence electrons in two, singly occupied e_g orbitals. However, these orbitals can be greatly affected by small distortions of the octahedral geometry.

The crystal field splitting parameter and the Ricah B parameter for β -Ni(OH)₂, assuming ideal octahedral geometry, have been reported: $\Delta_0 = 9000 \text{ cm}^{-1}$ and $B = 925 \text{ cm}^{-1.61}$ Yu et al. present reflectance data for precipitated Ni(OH)₂ and observe peaks at ~670 nm and ~385 nm,¹⁵ which correspond to the ν_2 (${}^{3}A_{2g} \rightarrow {}^{3}T_{1g}$) and ν_3 (${}^{3}A_{2g} \rightarrow {}^{3}T_{2g}$) transitions calculated for the β phase.⁶¹ Two additional peaks can be seen at ~740 nm and ~320 nm that are likely from α -Ni(OH)₂, although the study does not provide conclusive evidence regarding this assignment.¹⁵ Formally, the d–d transitions are forbidden by the Laporte rule,

Table 11. Comparison of Literature Values for the Vibrational Modes in α -Ni(OH)₂

	mode frequency (cm ⁻¹)	
our peak assignment	this study	literature frequency
Raman		
lattice mode	460	462-464, ³² 451-460, ³⁵ 460, ³⁶ 461 ³⁹
lattice mode	495	
2nd order lattice mode	~790	
2nd order lattice mode	~1075	
O–H bend, lattice OH	~1400	
O–H bend, layer H ₂ O	1590-1620	
O−H bend, free H₂O	1630	
O–H stretch, free H_2O	~3240, ~3380, 3515	
O–H stretch, lattice OH/layer H ₂ O	3590, 3620, 3642–3650	3625, ³¹ 3650–3653, ³² 3657, ³⁵ 3656, ³⁶ 3637 ³⁹
Infrared		
lattice mode		400, ²¹ 380 ²⁴
lattice mode		461, ¹ 474, ⁴ 460, ²¹ 480, ²⁴ 480 ²⁶
lattice mode	≤650-670	662, ¹ 645, ⁴ 625, ²¹ 650, ²⁴ 650–655 ²⁶
combination lattice mode	835	
combination lattice mode	1055	1055 ¹
O–H bend, lattice OH	1380-1390	1400^{1}
O–H bend, lattice OH	1480-1490	1540 ¹
O−H bend, layer H₂O	1600	1640, ¹ 1650, ¹⁴ 1650, ²¹ 1600, ²⁴ 1615–1620 ²⁶
$O-H$ bend, free H_2O	1630	
O–H stretch, free H_2O	3220/3240, 3380/3390, 3500/3515	~3400, ⁴ 3350, ¹⁴ 3350, ²¹ ~3500, ²⁴ ~3500, ²⁶ 3450 ⁴²
O–H stretch, lattice OH/layer H ₂ O	3570, 3600–3640	~3600, ⁴ ~3600 ²⁴

whereas such transitions are allowed if the crystal becomes asymmetric, i.e., if crystal disorder increases.⁶² An additional transition is observed above 33 000 cm⁻¹ (below 300 nm)¹⁵ that, based on its intensity, is from a charge transfer transition.

In the case of α -Ni(OH)₂ fluorescence is not observed for sample α_3 , which signifies that α -Ni(OH)₂ is not inherently fluorescent and suggests that the fluorescent background in sample α_2 may arise from disorder or changes in the electronic structure due to foreign Cl⁻ anions. In the case of β -Ni(OH)₂, both samples are fluorescent; however, they are also both partly disordered. Therefore, it is unknown whether the material is inherently fluorescent or whether more crystalline samples do not show a fluorescent background signal.

3.10. Disorder. The current body of literature can be simplified by considering two phases, α and β , with varying types and degrees of disorder. Turbostratic, or randomly oriented layer, structures can occur for α -Ni(OH)₂ samples, where adjacent layers are separated by amorphous H₂O. Regarding β -Ni(OH)₂, there is one kinetically stable layer orientation that forms by stacking faults during precipitation and during aging of α to β , as observed by XRD. The anions in the Ni^{II} salt can incorporate into the α -phase, as observed by XPS, Raman, and IR spectroscopy. Different degrees of hydration can occur, which is observed by large H₂O bands in the Raman and IR spectra. Finally, other factors such as crystallite size, point defects, stacking faults, etc. are expected to affect the spectroscopic properties of the materials, such as whether secondorder transitions appear in Raman measurements. In the case of stacking fault disorder, only the XRD peaks associated with the c-axis broaden, whereas other types of disorder cause all the diffraction peaks to broaden equally. However, the strong internal O-H stretching modes and the first order lattice transitions provide no evidence of any true additional crystal phases other than α and β .

These results show that changes in activity between samples that had previously been ascribed to the presence of intermediate phases actually arises from various types and degrees of structural disorder. By simultaneously, but separately, monitoring disorder from the incorporation of foreign ions, hydration and crystal defects such as stacking faults, changes in the performance, and stability of catalysts and electrodes can be linked with their true origin. This will allow for more systematic and more directed development of improved materials for devices.

4. CONCLUSIONS

This work presents a unified and simplified re-examination of previous spectroscopic investigations and demonstrates that one need only consider two phases (α and β) of Ni(OH)₂. Highly crystalline samples of α -Ni(OH)₂ may be impossible to produce because its layered structure minimizes its free energy by adopting some degree of disorder. However, highly crystalline samples of β -Ni(OH)₂ can be prepared, although the structure can also show disorder. For the first time, all of the Raman and IR-active lattice modes of β -Ni(OH)₂ have been identified and assigned using factor group analysis.

Samples can range in disorder from (1) incorporation of foreign ions, (2) hydration, and (3) crystal defects including stacking faults. The presence of foreign ions can be measured from characteristic vibrational modes, e.g., nitrate and sulfate anions, and/or by elemental analysis methods, such as XPS, e.g., chloride anions. Vibrational modes provide more detail on subtle changes in chemical environment, e.g., nitrate was shown to occupy two distinct chemical environments in α -Ni(OH)₂. Hydration affects the intensity of water bands in Raman and IR spectra. Increased disorder from crystal defects can be observed by the increased intensities of higher order vibrational modes in Raman and IR spectra and from the strong fluorescent backgrounds in the Raman spectra. Mechanical stress within the crystal is measured by observing small shifts in frequency of the internal vibrational modes. For example, at various locations within the sample or between similar samples. Further, stacking faults in β -Ni(OH)₂ samples can be measured by the presence of additional O–H stretching modes at 3601 $\rm cm^{-1}$ and 3652 $\rm cm^{-1}$

in the Raman and IR spectra, respectively. This type of disorder broadens only the XRD peaks involving the *c*-axis, whereas other types of disorder and small crystallite effects broaden all of the diffraction peaks equally. By examining different types of disorder individually, future studies on applications of Ni(OH)₂ will be able to link desired behavior with its structural origin.

Whereas thin films of α - and β -Ni(OH)₂ are difficult to detect or differentiate by other techniques, such as XRD and XPS, the two phases are easily distinguished from their strong internal O–H stretching modes at 3581–3640 cm⁻¹. The value of this is clear: for example, a re-examination of previous work reveals evidence of mixed samples from IR spectroscopy where XRD only detected the β -phase.¹ Thus, Raman or IR spectroscopy can be used to rapidly identify Ni(OH)₂ films by measuring these vibrational modes, which act as a fingerprint for pure and mixed samples (Figure 11).

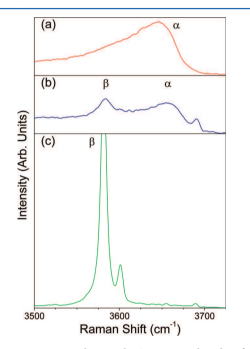


Figure 11. Raman spectra showing the O–H internal modes of samples (a) α_3 , (b) α/β , and (c) β_3 . All spectra are plotted on the same intensity scale.

This study provides the groundwork to determine which $Ni(OH)_2$ phase, or in the case of a mixture, the relative amount of each phase, is present at an electrode or catalyst surface and therefore to correlate desired electrochemical or catalytic behavior to a particular phase or degree of disorder. It also establishes a method to monitor change at the surface, i.e., the stability of an electrode or catalyst, during prolonged use or storage.

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Notes

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