FTIR INVESTIGATION OF THE EVOLUTION OF THE OCTAHEDRAL SHEET OF KAOLINITE-SMECTITE WITH PROGRESSIVE KAOLINIZATION

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Abstract—Twenty two samples were studied to investigate the nature and evolution mechanism of mixed-
layer kaolinite-smectite (K-S). We examined the \leq 2 µm or \leq 0.2 µm fraction of K-S formed by hydrothermal and hypergenic alteration of volcanic material. The samples are from three localities: 20 specimens from a Tortonian clay deposit in Almería, Spain; one specimen from weathered Eocene volcanic ash from the Yucatan Peninsula, Mexico; and one sample from a weathered Jurassic bentonite from Northamptonshire, England. The samples were studied using chemical analysis, X-ray diffraction (XRD) and Fourier transform infrared spectroscopy (FTIR). The XRD patterns of the oriented, glycolated mounts were modeled using NEWMOD and the proportion of smectite and kaolinite layers was determined, ranging between 0 and 80% kaolinite. The analysis of the OH-stretching region of the FTIR spectra at different temperatures $(180-550^{\circ}\text{C})$ showed the progressive dehydroxylation of kaolinite domains and, perhaps, of smectite domains, but no detailed information could be obtained about the sequential OH loss
in different cation environments. The abundance and short-range ordering of the octahedral cations were
studied using performance in different cation environments. The abundance and short-range ordering of the octahedral cations were studied using the OH-bending bands. The chemical and FTIR-estimated octahedral cation abundances were broadly similar. Aluminum showed a tendency to mix with Fe and Mg rather than to form AlAl pairs. Al-for-Mg substitution accompanying kaolinization was evident from the increase in AlAl pairs and decrease in AlMg pairs. Iron is retained in the structure. No other octahedral cation rearrangement was observed. The intensity of the 750 cm^{-1} band, assigned to translational vibrations of external OH groups in observed. The intensity of the 750 cm τ band, assigned to translational vibrations of external OH groups in
a kaolinitic environment, was quantified and modeled in relation to kaolinite layer proportion. The
chemical da chemical data show that there are residual interlayer cations in kaolinite domains, which in accordance with the model mentioned above, disturb external OH-translation vibrations. These results indicate the persistence of certain chemical and structural smectite features in kaolinite domains and thus support a smectite kaolinization process via a solid-state transformation. This confirms previous XRD, thermal, chemical and NMR analyses of the same sample set.

Key Words-Cation Distribution, FTIR, Kaolinite-smectite. Key WordsÐCation Distribution, FTIR, Kaolinite-smectite.

INTRODUCTION
The present contribution follows our recent study of mixed-layer kaolinite-smectite (K-S) (Dudek et al., 2006), which integrates the results of XRD, thermogravimetry (TG), chemical analysis and 29 Si nuclear magnetic resonance (NMR) to assess the structure and the mechanism of formation and transformation of K-S samples from both hydrothermal and hypergenic settings. We observed non-linear relations between the proportion of kaolinite and smectite layers measured by XRD and other parameters describing different aspects of the structure and chemical composition of $K-S$, such as percent weight loss due to the dehydroxylation of the octahedral sheet, position of the 06 XRD peak, the area of the kaolinite ²⁹Si NMR peak, MgO and $Fe₂O₃$ contents, and cation exchange capacity (CEC). Analysis of these trends led us to the conclusion that smectite kaolinization is a solid-state transformation that smective includes the solid-state transformation that proceeds through several non-simultaneous stages. The

progressive removal of tetrahedra and the increasing abundance of OH groups in the octahedral sheet lead to the development of kaolinite-like patches or domains within smectite layers. However, the development of the patches and their collapse to \sim 7 Å (as measured by XRD) do not progress simultaneously, because the collapse happens only when the patches acquire sufficient size. Also, other chemical and structural changes that occur during smectite-to-kaolinite transformation, such as Al-for-Mg replacement in the octahedral sheet and Si-for-Al in the tetrahedral sheet, and the resulting removal of interlayer cations, are not immediately linked to the collapse of layers to \sim 7 Å. We also observed that Fe is either retained in the structure of the newly formed kaolinite or released at a much slower rate than that of the transformation.

In the present paper we describe the results of FTIR applied to the same set of samples. The chief advantage of FTIR is that it gives a direct insight into the cationic environments of the OH groups and allows their qualitative and quantitative assessment.

Previous FTIR studies of K-S provided general qualitative information on the composition of K-S, using FTIR as a tool to detect smectite and kaolinite using FTIR as a tool to detect similarly and interesting by tracing the progressive appearance of bands char-

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acteristic of these two phases in the OH-stretching region (Churchman *et al.*, 1994; Srodon, 1980) or both OH-stretching and -bending regions (Delvaux et al., 1989, 1990). They emphasized the importance of the 3700 cm^{-1} OH-stretching vibrations to detect small amounts of kaolinite. However, no attempt was made to extract any further structural information on mixed layering. Madejová et al. (2002) performed a quantitative FTIR study of kaolinite dispersed in fine fractions of bentonite and found that the limit of detection of kaolinite is as low as 0.5 mass%. Churchman et al. (1994) and Delvaux et al. (1990) also discussed the FTIR signatures that help to differentiate between kaolinite and halloysite in mixed-layer sequences with kaolinie and halloysite in mixed-layer sequences with smectite.
In the present study we apply a method of FTIR

spectrum decomposition into individual metal-OH-metal bending bands, which allows relatively accurate quantification of cation pairs in the structure and assessment of cation ordering. The FTIR results show a lack of cation rearrangement during the reaction and thus contribute additional evidence for a solid-state transformation as additional evidence for a solid-state transformation as $t_{\rm eff}$ mechanism of smectra mechanism of smectra mechanism of small μ

MATERIALS AND METHODS

The samples investigated in this study represent both
hydrothermal and hypergenic alteration of volcanic material. Twenty samples are from a Tortonian clay deposit (Almería, Spain; Linares, 1985), one (RS49) is from a weathered Eocene volcanic ash (Yucatan, Mexico; Schultz et al., 1971), and one (CWP-73) from a weathered Jurassic bentonite (Northamptonshire, England; Hillier et al., 2002). The two last samples widened the range of K-S composition. The specimens from England and Mexico were pre-treated to remove amorphous Fe phases by the ammonium oxalate (Smith, 1994) and the dithionite-citrate-bicarbonate (Jackson, 1975) methods, respectively. The \leq μ m particle size fraction (0.2μ m for sample RS49) was separated from the bulk samples by dispersion in distilled water and sedimentation according to Stoke's law. The clay
separates contain virtually only K_S : trace amounts of separates contain virtually only K-S; trace amounts of quartz were found in four or them. Aliquots of the separated fraction were Li- and Ca-saturated in the exchangeable cation sites by dispersion in 3 M LiCl and 0.1 M CaCl₂ solutions, respectively. The solid/liquid ratio was $100 \text{ mg}/10 \text{ mL}$ and the samples were exchanged three to four times.

The XRD analysis was performed at the Natural History Museum (London, UK) using a Philips PW1050 diffractometer operating under the following conditions: 42 kV, 42 mA, Cu tube, graphite secondary monochromator, 1° divergence slit and 0.1 mm receiving slit. The Li-saturated samples were analyzed in oriented mounts prepared by sedimentation on glass slides (2.5 mg/cm^2) prepared by sedimentation on glass slides (2.5 mg/cm)
and recorded in air dried and glycolated forms (overand recorded in air-dried and σ

night saturation in ethylene glycol atmosphere at 60° C) from 3 to 30°2 θ , at 0.05°2 θ /step and 10 s counting time. The proportion of kaolinite and smectite layers in K-S (percent kaolinite layers, $\%$ K) was calculated by fitting the experimental patterns of the ethylene glycol-saturated samples with those calculated with the NEWMOD computer program (Reynolds and Reynolds, 1996).

Chemical analyses, including CEC, of the Los Trancos specimens are from Reyes et al. (1978) and Cuadros et al. (1994). These analyses were performed before cation exchange. The Mexico and England specimens were analyzed at the Natural History Museum, London, using an ICP-AES Varian Vista Pro apparatus after dilution in LiBO₂ (1:5), fusion and dissolution in $HNO₃$. These two samples were analyzed after cation exchange.

The TG analysis was performed at the Istituto di Metodologie per l'Analisi Ambientale, CNR, Potenza,
Italy using a Seiko Exstar 6000 apparatus. The samples Italy, using a Seiko Exstar 6000 apparatus. The samples were heated in Al_2O_3 crucibles from 20 to 1050°C at 10°C/min in a N_2 flow. Sample weights ranged from 9 to 15 mg. The dehydroxylation weight loss was measured from the TG curves, after discrimination from hydration water loss. For our samples, the temperature of the transition between dehydration and dehydroxylation ranges from \approx 260 to 400 °C.

We performed two types of FTIR experiments using a Perkin Elmer FTIR system at the Natural History Museum. In the first type, selected samples (representing a range of $\%$ K values) were analyzed in the Perkin Elmer AutoImage FTIR microscope equipped with a liquid-nitrogen-cooled mercury cadmium telluride (MCT) detector and a Linkam FTIR heating and cooling stage. A small amount of sample was placed on a ZnSe window, directly on the piece containing a thermopar that controls the temperature. The stage is closed and the IR beam enters and exits it through ZnSe windows. We analyzed single clay particles, thin and large enough to provide a reliable spectrum. The spectra were recorded from the same clay particle at ambient temperature and after heating to the following temperatures: $180-300^{\circ}$ C, 300–350°C, 420–430°C, 500°C and 550°C. The wide range of the lowest temperature (*i.e.* from 180 to 300°C) range of the lowest temperature (*i.e.* from 180 to 300°C) is due to the variable hydration-water content (lowest for the most kaolinitic sample) and ease of water removal (greater for kaolinite-rich K-S). For each sample is due to the variable hydration-water content (lowest for the most kaolinitic sample) and ease of water removal (greater for kaolinite-rich K-S). For each sample, 50 transmittance mode with a resolution of 4 cm^{-1} . The transmittance mode with a resolution of 4 cm \cdot . The
original interest in the heating experiments was based on
the fact that dehydroxylation in kaolinite occurs at lower
temperatures than in montmorillonite. We expected, the fact that dehydroxylation in kaolinite occurs at lower temperatures than in montmorillonite. We expected, therefore, to be able to identify the octahedral composition of smectite and kaolinite domains in K-S through monitoring the gradual loss of the OH groups related to the different cations with increasing temperature. However, we encountered problems that made the However, we encountered problems that made the attempt unduccessful . The OH-bending region of the OH-bending region of the σ

spectra from the heated samples was deformed as the SiO band at $\sim 1000 \text{ cm}^{-1}$ widened considerably and SIO band at \sim 1000 cm widened considerably and
enveloped the OH bands which thus became almost
imperceptible. The OH-stretching region of the spectra
was noisy and did not show sufficient detail. The enveloped the OH bands which thus became almost
imperceptible. The OH-stretching region of the spectra imperceptible. The OH-stretching region of the spectral hydration water band that overlaps smectite bands in the OH-stretching region remained up to relatively high temperatures and also complicated the analysis. However, we were able to investigate general changes. in the OH-stretching region during the consecutive stages of heating. This was done by subtracting the spectra recorded at higher temperatures from those recorded at the lowest temperature (excluding ambient temperature because the hydration water band was relatively prominent). The product of the subtraction was then decomposed into several bands that correspond to the approximate positions of kaolinite bands $(\sim 3700 \text{ cm}^{-1}$ and $\sim 3660 - 3640 \text{ cm}^{-1})$ and smeetite (~3700 cm $^{-1}$ and ~3660–3640 cm $^{-1}$) and smectite
bands (~3660–3640 cm⁻¹). The decomposition was
performed by curve-fitting, using the computer program
Galactic Grams/AI (version 7.01). The areas of the bands
in the bands (~3660–3640 cm
performed by curve-fitting,
Galactic Grams/AI (version
in the subtracted spectra we
the areas of the individu
recorded at the lowest temp performed by curve-fitting, using the computer program Galactic Grams/AI (version 7.01). The areas of the bands $\frac{1}{\pi}$ (version $\frac{1}{\pi}$). The areas of the bands of the ba the areas of the individual bands from the spectra recorded at the lowest temperature.

In the second type of experiments, all samples were analyzed as KBr pellets (sample/KBr ratio of: 1 mg/200 mg) using a Perkin Elmer Spectrum One workbench equipped with a deuterated triglycine sulphate (DTGS) detector with a CsI beamsplitter. Spectra were recorded in the range from 4000 cm⁻¹ to 300 cm⁻¹ with a 5 cm⁻¹ resolution, in transmission mode. The region of the OHbending vibrations (940–550 cm⁻¹) was selected for a bending vibrations (940–550 cm
detailed analysis because it does no
with water hydration bands and be
better resolved. First, a baseline was
in Figure 1. The baseline was calcu
polynomial function after fixing its detailed analysis because it does not show interference ariance analysis because it does not because the bands are better resolved. First, a baseline was subtracted as shown in Figure 1. The baseline was calculated using a quintic polynomial function after fixing its value at both sides of the part of the spectrum to be analyzed. There is uncertainty about the accuracy of the position of the baseline. The accuracy of this procedure and the overall baseline. The accuracy of this procedure and the overall \mathcal{A}

Figure 1. Baseline (dashed) subtracted from the OH-bending region of the original spectrum before its deconvolution and region of the original spectrum before its deconvolution and
decomposition into single metal.OH.metal bands decomposition into single metal-OH-metal bands.

were then deconvoluted in order to find the band maxima
and finally decomposed by curve-fitting into the individual metal-OH-metal bands (metal = Al, Mg, Fe) as Gaussian-shaped functions. The intensities of the bands were normalized with respect to the sum of the areas of all OH-bending bands to allow comparison areas of all OH-bending bands to allow comparison
between complex between samples.

RESULTS AND DISCUSSION RESULTS AND DISCUSSION

XRD measurements of kaolinite:smectite layer

Table 1 shows the NEWMOD-calculated compositions of K-S samples and the parameters used for the calculations. Selected examples of the XRD experimental and modeled patterns are presented in Figure 2. The $\%$ K indicates the percentage of kaolinite layer content in K-S. A satisfactory fit for the peak positions and widths was obtained for all samples. However, we encountered problems with the fitting of the background, encountered problems with the thing of the catagoritms, ϵ is the low-angle region. For the most the most the most that ϵ

Figure 2. Selected XRD patterns of the oriented, ethylene
glycol-solvated K-S samples and their NEWMOD-modeled traces (black: experimental, gray: calculated). Peak positions in A. (a) smectite (TR-4); (b) 16% K (LT-9/1); (c) 32% K (TR-A. (a) smeetite (TR-4), (b) 10% K (LT-9/1), (c) 32% K (TR-
125): (d) 40% K (TR-121): (e) 80% K (RS40) $(1, 0)$; (e) $(1, 1)$; (e) $(2, 1)$; (e) $(2, 1)$; (e) $(2, 1)$; (e) $(3, 1)$; (e) $(4, 1)$; (e)

Sample	Total %K	Phase	$\%K$ in each phase	Wt.% of phase	$d_{\rm K}$ (Å)	N_{ave}	$\cal N$
TR-4	$\boldsymbol{0}$	smectite		100		$2.5\,$	$1 - 10$
TR-146	$\boldsymbol{0}$	smectite		100		2.5	$1 - 10$
$T-21$	7	$K-S$	$\overline{7}$	100	7.4	2.5	$1 - 10$
TR-3	$\overline{7}$	$K-S$	$\sqrt{ }$	100	7.4	2.5	$1 - 10$
$T-16$	10	$K-S$	10	100	7.4	2.5	$1 - 10$
TR-2	13	$K-S$	13	100	7.4	2.5	$1 - 10$
$LT-9/1$	16	$K-S$	10	90	7.4	2.5	$1 - 10$
		$K-S$	60	10	7.15	2.5	$1 - 10$
$T-22$	25	$K-S$	13	70	7.4	2.5	$1 - 10$
		$K-S$	50	$30\,$	7.4	2.5	$1 - 10$
$LT-10/6$	32	$K-S$	20	60	7.4	2.5	$1 - 10$
		$K-S$	60	30	7.15	2.5	$1 - 10$
		smectite	—	10		2.5	$1 - 10$
TR-119	32	$K-S$	20	80	7.4	2.5	$1 - 10$
		$K-S$	70	$20\,$	7.15	2.5	$1 - 14$
TR-125	32	$K-S$	15	65	7.4	2.5	$1 - 10$
		$K-S$	60	35	7.15	2.5	$1 - 10$
$LT-9/3$	33	$K-S$	25	75	7.4	2.5	$1 - 10$
		$K-S$	60	20	7.15	2.5	$1 - 10$
TR-115	34	$K-S$	25	60	7.4	2.5	$1 - 10$
		$K-S$	60	30	7.15	2.5	$1 - 10$
		smectite	$\overline{}$	10	$\overline{}$	2.5	$1 - 10$
$T-18$	37	$K-S$	20	70	7.4	2.5	$1 - 10$
		K-S	70	30	7.15	2.5	$1 - 14$
TR-117	38	$K-S$	38	100	7.4	2.5	$1 - 10$
$LT-9/2$	39	$K-S$	25	55	7.4	2.5	$1 - 10$
		$K-S$	60	35	7.15	2.5	$1 - 10$
		smectite		10		2.5	$1 - 10$
$T-20$	47	K-S	30	70	7.4	2.5	$1 - 10$
		$K-S$	80	30	7.15	2.5	$1 - 14$
TR-121	49	$K-S$	40	85	7.4	2.5	$1 - 10$
		$K-S$	90	15	7.15	2.5	$1 - 14$
$LT-2$	55	$K-S$	40	75	7.4	2.5	$1 - 10$
		$K-S$	90	25	7.15	2.5	$1 - 14$
$LT-10/1$	61	$K-S$	30	55	7.4	2.5	$1 - 10$
		$K-S$	90	42	7.15	2.5	$1 - 14$
		kaolinite	100	3	7.12	$10\,$	$1 - 55$
RS49	80	$K-S$	80	100	7.15	3	$1 - 14$
$CWP-73$	85	$K-S$	85	90	7.15	2.5	$1 - 14$
		Illite		10	$\overline{}$	4	$1 - 14$

 Γ 1. The NEWMOD-calculated compositions of the K-S samples (clay fraction) and the parameters (clay fraction) and

 $1-14$
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ample coherent scattering domains. Other parameters used for modeling: d value of ethylene glycol-solvated smectite layers: 16.9 Å for RS49 and CWP-73, 16.7 Å for all the other samples; d value of illite (sample smectite layers. 16.9 A 161 KS49 and CW1-75, 16.7 A 161 and the other samples, d value of filtre (sample
CWP-73): 9.94 Å: ordering: $R = 0$ excent for CWP-73 and RS49 where $R = 1$: $\sigma^* = 30$ CWP-73). 9.94 A, ordering: $R = 0$ except for CWP-73 and RS49 where $R = 1$, $Q = 30$.

 k and consider $\frac{1}{2}$ and $\frac{1}{2}$ misite in the intensity of the peak around 20 20 , which
we were not able to solve we were not able to solve.
For eight samples, there was a good fit of the

experimental and theoretical patterns using one phase (smectite or K-S, Figure 2a,e). For the other 12 samples, two or three phases were needed, including K-S of different composition (kaolinite-poor: $\leq 50\%$ K, and kaolinite-rich: $>50\%$ K), smectite, and, in two cases, kaolinite and illite (Figure 2b-d). K-S with $\leq 50\%$ K was a dominant component of the mixtures, comprising $55-90$ wt.% (Table 1). This heterogeneity may be a dominant component of the mixtures, componing $\frac{1}{2}$

related to the differential transformation of smectite to kaolinite within a confined space. However, the model of phase mixing applied to simulate this heterogeneity. probably does not represent the exact composition of the corresponding samples.

Some of the calculation parameters for kaolinite-poor and kaolinite-rich K-S were different. The 7.4 \AA d spacing of the kaolinitic layers in K-S with $\leq 50\%$ K may suggest that they are non-expanding halloysite rather than kaolinite (e.g. Brown and Brindley, 1980). We were not able to differentiate between kaolinite and halloysite not able to differentiate between the continuous and hallows of $\sum_{i=1}^{n}$

Theng *et al.*, 1984), based on halloysite layer swelling, also cause the expansion of smectite layers, and therefore are not suitable for K-S samples. However, Brindley (1980) suggested that halloysite is a highly disordered form of kaolinite and, in K-S with low kaolinite content, disordered kaolinite layers are more likely to be present than well crystallized kaolinite layers. As it is not possible to discriminate between the two, for simplicity we refer to the 7.4 \AA component as kaolinite, in line with many previous studies describing K-S.

The values for the goniometer radius (17.3 cm) and sample length (2.5 cm) from our diffractometer were used in the NEWMOD calculations. The Fe content in smectite was taken from the chemical data, recalculated for the half unit-cell of a 2:1 dioctahedral clay structure. The Fe content of our samples ranges from 0.03 to $0.2/O_{10}(OH)_{2}$. We assigned all Fe in a given sample to smectite layers, although this appears not to be the case. We found, however, that the patterns calculated using different Fe content (within the range of Fe values in our samples) in smectite and 7 Å layers are identical. Kaolinite layers were calculated without Fe. Kaolinite layers were calculated without Fe.

FTIR data
OH-stretching region. Figure 3 shows the stretching region $(3800-3400 \text{ cm}^{-1})$ of the FTIR spectra (workregion (3800–3400 cm) of the FTIR spectra (work-
bench experiments) for selected samples arranged
according to increasing % K. As expected, there is a $\frac{1}{2}$ becoming to increasing $\frac{0}{6}$ K As expected there is a according to increasing β , there is an expected, there is a

Figure 3. The OH-stretching region of selected samples (TR-4, T-16, TR-119, T-20, LT-2, RS49). The figure shows progressive development of the \sim 3700 cm⁻¹ band and sharpening of the \sim 3620 cm⁻¹ band with increasing kaolinite content in K-S \approx 3620 cm \cdot band with increasing kaolinite content in K-S
(expressed as % K in the figures at right of the plot). \mathcal{C} is a figure set of the figures at right of the plot \mathcal{C}

progressive growth of the 3700 cm⁻¹ band, correspond-
ing to AlAlOH-stretching vibrations in kaolinite (e.g. Russell and Fraser, 1994) with increasing degree of kaolinization. It should be noted that this band was not observed in the samples that are characterized by XRD as pure smectite (TR-4 and TR-146, Table 1). However, according to our TG results, these samples already. contain kaolinite domains. Although the 3700 cm^{-1} contain kaolinite domains. Although the 3700 cm
region is very sensitive to the presence of kaolinite in
mineral mixtures (Madejová *et al.*, 2002), kaolinite
crystals in a physical mixture with smectite are not region is very sensitive to the presence of th crystals in a physical mixture with smectite are not equivalent to kaolinite-like domains within smectite layers in K-S. Kaolinite-like domains are likely to be affected by the surrounding smectite structure and their 3700 cm^{-1} band can be expected to be less intense than in well crystallized kaolinite. Thus, a number of AlAlOH groups in a kaolinite-like domain in mixed-layer K-S will produce a band of lower intensity than the same number of AlAlOH groups in a kaolinite crystal. It seems, therefore, that TG is more sensitive than XRD and FTIR for detecting kaolinite in K-S.

Churchman et al. (1994), Delvaux et al. (1990) and Watanabe et al. (1992) inferred that the several soil samples they studied had halloysite, rather than kaolinite, interstratified with smectite because they found no band at 3650 cm⁻¹. In our samples, this band only band at 3650 cm \therefore In our samples, this band only
appears in those containing $\geq 80\%$ K (Figure 3). In our
view, this may be due to a constant increase in
crystalline order in the kaolinitic structure with increasview, this may be due to a constant increase in crystalline order in the kaolinitic structure with increasing kaolinization, in which the structure progresses from a halloysite-like stage to a disordered kaolinite-like stage, as observed by IR (Russell and Fraser, 1994). Also, the 3650 cm^{-1} band could be masked in the less kaolinitic specimens by the smectite band.
Heating-experiment analysis of the OH-stretching kaolinitic specimens by the smectite band.

region. The heating experiments were conducted using the microscope on seven samples with a $\%$ K range of 0 to 85 (TR-4, TR-119, T-20, TR-121, LT-2, RS49, CWP-73, see Table 1). Figure 4 shows the OH-stretching region of the spectra recorded at the consecutive stages of heating for a sample of intermediate $\%$ K content. As expected, due to progressive dehydroxylation, there is a gradual loss of absorption intensity as heating progresses, and the absorption loss seems greater in the \sim 3700-3600 cm⁻¹ region. There is also decreased absorption in the water-hydroxyl region (below 3600 cm^{-1}). In order to evaluate the loss of absorption 3600 cm 3). In order to evaluate the loss of absorption
corresponding to structural hydroxyls we subtracted the
spectra recorded at the higher temperatures from that run
at the lowest temperature for all samples subjected spectra recorded at the higher temperatures from that run at the lowest temperature for all samples subjected to the heating experiments. Figure 5 shows two subtracted spectra and their decomposition for sample T-20 (the same as in Figure 4). At 420° C, the absorption loss occurs at \sim 3700 cm⁻¹, while at 500°C it occurs at both occurs at \approx 3700 cm
 \sim 3700 cm⁻¹ and \sim 366
In some of the spect: ~3700 cm⁻¹ and ~3660-3640 cm⁻¹ (thick gray lines). ~3700 cm \degree and ~3660-3640 cm \degree (thick gray lines).
In some of the spectra, the ~3660-3640 cm⁻¹ region In some of the spectra, the \sim 3660 $-$ 3640 cm \sim region

 $F_{\text{e,} \text{c,} \text{d}}$ at three subsequent stages of heating $F_{\text{e,} \text{d}}$ recorded at three subsequent stages of heating.

could not be modeled precisely due to the influence of the hydration water bands (thin black lines). Figure 5 is a good example of the results of this procedure: the intensity loss at low temperature occurred only at 3700 cm^{-1} or was higher than that at $3660 - 3640$ cm⁻¹, but at higher temperature the relative 3660–3640 cm $^{-}$, but at nigher temperature the relative
amount of intensity loss at 3660–3640 cm⁻¹ increased.
This is due to the fact that kaolinite dehydroxylates at
lower temperature than montmorillonite. Indeed, ka This is due to the fact that kaolinite dehydroxylates at lower temperature than montmorillonite. Indeed, kaolinite has maxima at 3700 and 3620 cm⁻¹ while the smectite maximum is at $\sim 3620 \text{ cm}^{-1}$. lower temperature than montmorillonite. Indeed, kaolinite has maxima at 3700 and 3620 cm^{-1} while the

ite has maxima at 3700 and 3620 cm⁻¹.
Since the assessed the absorption loss at the subsequent
temperatures by representing the areas of the bands at smectite maximum is at ~3620 cm
We assessed the absorption loss
temperatures by representing the are
~3700 cm⁻¹ and at ~3600-3640 cm⁻¹ We assessed the absorption loss at the subsequent We assessed the absorption loss at the subsequent $\approx 3700 \text{ cm}^{-1}$ and at $\approx 3600 - 3640 \text{ cm}^{-1}$ in the subtracted ~3700 cm \sim and at ~3600–3640 cm \sim in the subtracted spectra as a fraction of the areas of the individual bands at the lowest temperature. Figure 6 is the result of these calculations and illustrates the evolution of spectra as a fraction of the areas of the individual bands at the lowest temperature. Figure 6 is the result of these calculations and illustrates the evolution of the absorption loss at the consecutive stages of heating for the two regions \sim 3700 cm \sim and \sim 3660–3640 cm
the absorption loss increases with in
The fraction of absorption loss for
always greater in the 3700 cm⁻¹ regi
heating experiments show mainly idroxylation of kaolinite layers in F the absorption loss increases with increasing temperature. The fraction of absorption loss for each temperature is T_{1} absorption of absorption T_{2} for each temperature is absorption of the state in the 3700 cm⁻¹ region indicating that the always greater in the 3700 cm Tregion, indicating that the heating experiments show mainly the progressive dehy-
droxylation of kaolinite layers in K-S. However, partial
smectite dehydroxylation is probably also happening heating experiments show mainly the progressive dehy-
droxylation of kaolinite layers in K-S. However, partial arring mirrie to antimate any result of the services, partial higher temperatures. The combination of the absorption loss mgers temperatures. The communities of the absorption loss

Figure 5. Results of the subtraction of the spectra recorded at 420°C and 500°C from that recorded at 230°C for sample T-20. The spectra are decomposed into several bands: the thick gray lines, labeled with their wavenumbe $(\sim 3700 \text{ cm}^{-1})$ and kaolinite plus smectite ($\sim 3660 \text{ cm}^{-1}$); the thin black lines correspond to hydration water.

Figure 6. Loss of absorption in the OH-stretching region at several temperatures corresponding to the areas in the subseveral temperatures corresponding to the areas in the subtracted and decomposed spectra (as shown in Figure 5).

spread of the band position values for the \sim 3660–3640 cm⁻¹ region. Another cause is the fact that this region is affected by overlap with the water-hydroxyl
band. This overlap made it impossible to obtain more
detailed information about K-S dehydroxylation.
OH-bending region. Figure 7 shows an example of the band. This overlap made it impossible to obtain more detailed information about K-S dehydroxylation. detailed information about K-S dehydroxylation.

OH-bending region $(960-800 \text{ cm}^{-1})$ of a FTIR spectrum OH-bending region (960–800 cm
(sample TR-117) decomposed into
metal bands after baseline subtraction
750 and 790 cm⁻¹ correspond
molecular interaction and will be
remaining bands (thin black lines (sample TR-117) decomposed into individual metal-OH-(sample Trees of the second into individual metal-out of the second into the second individual metal-OH-750 and 790 cm⁻¹ correspond to another type of 750 and 790 cm correspond to another type or
molecular interaction and will be addressed later. The
remaining bands (thin black lines) are not used here
although they are necessary in the curve-fitting calculamolecular interaction and the contraction and the contraction of the addressed later. remaining bands (the black lines) are not used here

tion. In all studied samples we detected the following OHuniversally accepted scheme for smectite and kaolinite (e.g. Russell and Fraser, 1994): 938 cm⁻¹ and 916 cm⁻¹: (e.g. Russell and Fraser, 1994): 938 cm π and 916 cm
AlAlOH; 870–890 cm⁻¹: AlFeOH; 835 cm⁻¹ and
845 cm⁻¹: AlMgOH; 815 cm⁻¹: FeFeOH; 780 cm⁻¹
MgFeOH. There is an additional band at ~895 cm⁻¹ tha AlAlOH; 870-890 cm⁻¹: AlFeOH; 835 cm⁻¹ and ATAIOH; 870–890 cm \cdot : AIFEOH; 835 cm \cdot and
845 cm⁻¹: AlMgOH; 815 cm⁻¹: FeFeOH; 780 cm⁻¹:
MgFeOH. There is an additional band at ~895 cm⁻¹ that
develops with increasing kaolinite content and was
assigned to A 845 cm \cdot : AlMgOH; 815 cm \cdot : FeFeOH; 780 cm \cdot :
MgFeOH. There is an additional band at ~895 cm⁻¹ that
develops with increasing kaolinite content and was
assigned to AlAlOH (Frost, 1998). develops with increasing kaolinite content and was assigned to AlAlOH (Frost, 1998).

In Figure 8 we compare the abundance of the three octahedral cations (Al, Mg and Fe) determined from the FTIR and chemical analysis in order to test the reliability of the baseline correction and decomposition technique of the FTIR spectra. The metal abundances are expressed in the form of a 2:1 structure unit-cell, even though this does not have a physical meaning due to the presence of kaolinite, which has a $1:1$ structure. (Therefore, the octahedral occupancy for samples with the highest kaolinite content, RS49 and CWP-73, is >4). However, the figures are only considered in their relative values for comparison between chemical and FTIR data. In the case of Mg, the chemical data were corrected for the presence of interlayer Mg. However, the chemical data include tetrahedral as well as octahedral Al, because it is impossible to separate them in K-S. The FTIR cation content was calculated from the areas of the individual bands assuming that all metal-OH-metal vibrations have the same absorptivity. Figure 8a shows that there is a relatively good agreement between the FTIR and chemical assessment of Al content. The typical differences in Al estimations between FTIR and chemical analysis are $0.007-0.62$ atoms, yielding 4 to 13% relative error. The agreement is good in spite of the fact that the chemical data include tetrahedral Al, which fact that the chemical data include tetrahedral along a model.

Figure 7. FTIR spectrum of sample $\frac{1}{2}$ and $\frac{1$ dotted and dashed lines), possible translational vibrations of external OH groups (thick gray lines) and others calculated to obtain a good fit (thin black lines).

 ϵ comparison of the octahedral Al, ϵ comparison of the octahedral Allexandral Allow a direct comparison between the plots chemical analysis and FTIR. The axes show the same range of values to allow a direct comparison between the plots.

Al is low as shown by the chemical analyses of the smectite end-member of the series. The agreement is also relatively good for Mg (Figure 7b), with slightly greater values for the chemical analysis. The typical differences range from 0.07 to 0.26 atoms, yielding relative errors in the range 8 to 47%. However, FTIR noticeably overestimates Fe abundance with respect to the chemical analysis (Figure 8c). The typical differences range from 0.26 to 0.57 Fe atoms, producing large relative errors from 27 to 493%. We believe that this is mainly due to the difficulties in the accurate quantification of the small FeFeOH band, as the other band contributing to Fe abundance, AlFeOH, does not seem to induce a consistent error in the quantification of Al. Another source of error may have been the selection of the baseline. It seems, therefore, that the applied method of the FTIR spectrum decomposition produced broadly reliable results, with the exception of the FeFeOH band, which is, however, usually very small. In view of this, we decided to use the FTIR data for further analysis.

Figure 9 shows the relative absorption intensities of Figure 1.1 Shows the relative absorption intensity absorption in the relative of the relative of the relative of t $t_{\rm H}$ and $t_{\rm H}$ the cation pairs as a and $t_{\rm H}$

function of % K. As expected, the AIAIOH and AIMgOH bands show the opposite trend with progressing kaolinization of smectite, AlAlOH absorption increases and AlMgOH absorption decreases (Figure 9a,b). The intensity of the AlFeOH band shows a slightly decreasing. trend with increasing $\%$ K (Figure 9c). These three bending vibrations (AlAlOH, AlMgOH and AlFeOH) are the most reliable because their bands are relatively large and well resolved. However, in order to obtain a more complete picture of Mg and Fe behavior during kaolinization, all bands associated with these cations should be considered. The two remaining bands identified in the samples, MgFeOH and FeFeOH are usually small and probably affected by larger relative error; they do not show any trend as kaolinization proceeds (Figure 9d,e). The addition of the FeMgOH and AlMgOH bands causes little change of the trend in Figure 9b. This trend clearly shows the continuous removal of Mg from the K-S structure during kaolinization of smectite. In the case of Fe, however, the addition of the MgFeOH and FeFeOH bands to the AlFeOH band does affect the trend in Figure 0c, causing increased $\frac{d}{dx}$ affect the trend in Figure 9c, causing increased

and the sum of all bands involving Fe (f) as a function of % K (% of kaolinite layers). R^2 values correspond to a 95% confidence level.

spread of the data points (there is no meaningful correlation in Figure 9f). This implies that Fe remains in the structure of the newly formed kaolinite or is removed very slowly as the transformation proceeds, in agreement with our previous conclusions based on chemical analysis. We believe that this is supported by the FTIR data even if the band decomposition method overestimates Fe abundance (Figure 8c), because the data point trend, or the lack of it, depends on the relative intensity values and not on their absolute values. The fact that Fe is retained in the kaolinite layers indicates. that kaolinization of smectite proceeds through the inheritance of the original structure.

Figure 10a shows the change of the relative intensity of the band at 750 cm⁻¹ with the % K increase in the of the band at 750 cm \cdot with the % K increase in the K-S samples. Farmer (1974) related this band to translational vibrations implicating the external OH groups in kaolinite. Bishop *et al.* (2002) assigned it to $\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1}{\sqrt{1-\frac{1$ groups in kaolinite. Bishop et al. (2002) assigned it to out-of-plane OH bending rather than translation, without specifying whether the corresponding OH groups are external. However, as this band is characteristic of kaolinite, it is likely that the OH groups involved are external. This band is coupled in kaolinite with that at 790-795 cm^{-1} (Russell and Fraser, 1994) (see 790–795 cm⁻¹ (Russell and Fraser, 1994) (see
Figure 7); for this analysis, however, we used only the
band at 750 cm⁻¹ because it appears relatively free from
overlap with other bands and it is easier to quantify. The
 $F_{\rm{1}}$ and at 750 cm^{-1} because it annears relatively free from band at 750 cm because it appears relatively free from
overlap with other bands and it is easier to quantify. The
tetrahedral Al-O-Si in-plane vibration band also appears
at the same position of 750 cm⁻¹ (Farmer, 1974; overlap with other bands and it is easier to quantify. The tetrahedral Al-O-Si in-plane vibration band also appears tetrahedral al-O-Si in-plane vibration band also appears
at the same nosition of 750 cm^{-1} (Earmer, 1074; Pussell and Fraser, 1984) but the intensity of this band is very small in smectite and non-existent in kaolinite due to the low or absent tetrahedral Al, and it has very little or no influence on the present analysis. The intensity of the 750 cm^{-1} band was normalized with respect to the sum of all OH-bending bands plus the 750 cm^{-1} band itself.
The plot in Figure 10a shows that the datapoint trend

of all OH-bending bands plus the 750 cm \cdot band itself.
The plot in Figure 10a shows that the datapoint trend
is best approximated by a curve with two inflection
points (thin black line: a polynomial regression of the is best approximated by a curve with two inflection $\frac{1}{\pi}$ is best approximated by a curve with two interestion of the experimental data). This trend has a general shape similar to the equation in Figure 10b, although similar to the equation in Figure 10b, although `stretched' in the diagonal direction. Figure 10b shows

the increase of OH groups in K-S with progressing kaolinization of smectite. The non-linear relation in this plot is interpreted as being due to the development of kaolinite-like patches that collapse to \sim 7 Å only when they are sufficiently large (see Introduction). The similarity between the trends in Figures 10 a and b suggests that the cause of the intensity increase of the 750 cm^{-1} band is related to the increase in the number of OH groups in K-S, which should be the case if the assignment by Farmer (1974) (and possibly Bishop et $al., 2002$) is correct. However, since the datapoint trend in Figure 10a does not correspond exactly to that in Figure 10b, we considered the possibility that the intensity of the 750 cm^{-1} band does not depend on the number of external OH groups only. This vibration of external OH groups could be affected by the presence of interlayer cations and water. Neither of the two exists in end-member kaolinite but K-S does have residual interlayer cations in \sim 7 Å layers, as shown in Figure 10c. If there were no such residual interlayer cations in K-S, Figure 10c would show a linear trend. The residual interlayer cations are probably partially hydrated. Thus, the external OH vibrations considered may very well be affected by the interlayer complex, and the resulting intensity may be diminished. If this is the case, Figure 10a is produced by a combination of the increase in OH groups (Figure 10b) and the decrease in the CEC (Figure 10c), where both effects cooperate in the intensity increase of the 750 cm^{-1} band.

the intensity increase of the 750 cm shand.
We tested this hypothesis by modeling the
trend in Figure 10a using two components,
sion equation from Figure 10b (equation We tested this hypothesis by modeling the datapoint sion equation from Figure $10b$ (equation 1) and the inverse of the regression equation from Figure $10c$ $equation$ 2). These equations were modified to make their intercepts equal to zero as no intensity of the investigated band was measured for $\%$ K = 0. The equation in the model is $y = a$ (eq. 1) + b (eq. 2), where a and b are two parameters fitted experimentally. The best fit is shown in Figure 10a (thick gray line) and corresponds to the values $a = 1.5$ and $b = 0.15$. The fit corresponds to the values a series and the fitter can \mathbf{q} is good up to 80% keeps and the non-department and then dependent of \mathbf{q}

Figure 10. Intensity of the 750 cm $^{-1}$ band (a); percent weight loss due to dehydroxylation of the octahedral sheet (1G analysis) (b); and CEC (c) as a function of % K in K-S (XRD measurements). The thick gray line in (from plots (b) and (c). The thin black line in (a) and the B^2 value correspond to a polynomial regression of the experimental data. from plots (b) and (c). The thin black line in (a) and the R² value correspond to a polynomial regression of the experimental data.
Filled gray square: ideal value for smectite: onen square: ideal value for kaolinite. A Filled gray square: ideal value for smectite; open square: ideal value for kaolinite. All R² values correspond to a 95% confidence
_{Jevel}

from the experimental regression. However, the fact that we only have two experimental values in the region above $~60\%$ kaolinite layers generates some uncertainty about the actual trends in the experimental datasets involved in the model. The values of the parameters a and b are very different because equation 1 covers a numerical range of values of 11, whereas equation 2 covers a range of 110, *i.e.* 10 times larger. Thus, the fact that a is exactly 10 times larger than b indicates that the two equations (or the two variables: OH number increase and CEC decrease) have a similar weight in shaping the behavior of the trend in Figure 10a. In our opinion, the assumed effects on the intensity of the 750 cm^{-1} band are very likely and the good fit obtained supports the validity of our interpretation. Interestingly, however, the OH in-plane bending vibrations $(938-780 \text{ cm}^{-1})$ do not OH in-plane bending vibrations (938–780 cm
show the effect of the presence of residual in
cations in kaolinite domains. This can best be
Figure 9a, where the data point trend is linear. T
two reasons for this: (1) in this show the effect of the presence of residual interlayer cations in kaolinite domains. This can best be seen in cations in the catter account of the settle of the seed in the seed in Figure 2.1 and the seed in Figure 3.1 and the seed in Figure 3.1 and the seed in Figure 3.1 and the seed in the set of the seed in the set of the set o two reasons for this: (1) in this case both external and internal OH groups are implicated, and not just the external ones, as in the case of the 750 cm^{-1} band; and (2) the energy and geometry of the OH in-plane bending vibrations are different from those of OH-translation or OH out-of-plane vibrations.

We also investigated the short-range cation ordering in the octahedral sheet by comparison of the FTIR and chemical data. The charts in Figure 11 plot the ratio of the abundance of a selected cation over the sum of all cations from the chemical analysis $vs.$ the intensity of a catter from the catalogue analysis versus intensity of a \mathcal{S} and selected cation divided by the sum of

the intensities of all bands involving this cation.
Figure 11a evaluates the distribution of the AlAl pairs with respect to AlMg and AlFe. The greater chemical than FTIR ratios indicate that Al tends to mix with Mg and Fe more than occur as AlAl pairs. However, this tendency decreases with increasing kaolinite content as the trend approaches the 45° line, which indicates random distribution of Al in the octahedral sheet. This is expected because as kaolinization proceeds, more Al is incorporated in the octahedral sheet replacing Mg. However, a random Al distribution is not achieved even for samples containing 80% kaolinite. Figures 11b and c show that Mg and Fe mix with Al to a great extent, confirming the result from Figure 11a. Finally, Mg and Fe are segregated from each other, as shown in Figure 11d, except in one case (sample CWP-73), in which they appear to mix to a large extent. The distribution of the FeFe pairs cannot be reliably assessed. due to the ambiguous quantification of the FeFeOH band. The results above are not affected by the tetrahedral Al (accounted for in the chemical but not in the FTIR data). which is only a small fraction of the total Al.

In conclusion, it appears that the only change in the octahedral cation ordering of K-S samples during the smectite-to-kaolinite reaction is an increase in the number of the AlAl pairs, as shown by a trend in Figure 11a. The lack of trend in Figures 11b,c and d indicates that Mg and Fe are not rearranged during the reaction. The increment of AlAl pairs results only from the loss of Mg with increasing kaolinization. The Al is the loss of Mg with increasing increasing the City of the Al is the Al not randomly distributed at 80% K possibly because F

Figure 11. Cation ordering in the octahedral sheet. The plots show the intensity ratio of some individual metal-OH-metal bands to the sum of several metal-OH-metal bands, vs. the ratio metal/Al+Mg+Fe from the chemical anal sum of several metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-OH-metal-
Sum of the corresponding metal-oh-metal-oh-metal-oh-metal-oh-metal-oh-metal-oh-metal-oh-metal-oh-metal-oh-met a random cation distribution; data above and below the line indicate and below the corresponding metal pair higher and lower
than random respectively than random, respectively.

is preserved in the structure and AlFe pairs are still very abundant, as indicated in Figure 11c. These observations strongly suggest a solid-state transformation as the strongly suggest a solid-state transformation as the transformation of ϵ mechanism of kaolinization of smectite in these samples.

(1) The evolution of the intensity of the 750 cm
d in K-S, possibly corresponding to vibrations
blicating external (kaolinite-type) OH groups, with
gressing kaolinization is controlled by the number of band in E-S, possibly corresponding to vibration progressing kaolinization is controlled by the number of external OH groups in kaolinite domains, as expected, but also by residual interlayer cations (possibly hydrated). If the assignment of the band is correct, this result indicates that residual layer charge is preserved in the kaolinite domains.

 (2) During the smectite-to-kaolinite reaction, the intensities of the FTIR bands involving Mg decreases sharply whereas the intensity of the bands related to Fe do not show any trend. This is in agreement with the chemical data showing Mg loss and Fe retention in K-S with proceeding kaolinization.

(3) Analysis of short-range ordering in K-S octahedral sites indicates that atoms are not rearranged during smectite-to-kaolinite conversion, except that there is a decrease in AlMg pairs and an increase in AlAl pairs produced by Al-for-Mg substitution.

(4) These conclusions point to solid-state transformation as the most likely mechanism of kaolinization of smectite, in agreement with previous studies based on smective, in agreement with previous studies studies based \mathcal{S} , \mathcal{S} , and \mathcal{S} and \mathcal{S} and \mathcal{S} and \mathcal{S} and \mathcal{S}

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