Sodium hydroxo titanate nanowires with NaTi$_2$O$_4$(OH) formulation are fabricated by chimie-douce from TiO$_2$ powders in a hot concentrated base solution. In the present work, we investigate, in detail, the structure and morphology of the titanate, acid-exchanged materials, and phases formed during dehydration of these compounds; we then discuss the relationship between the initial titanate structure and TiO$_2$ formation. XRD, ED, HRTEM, EDS, density measurements, and TG/MS results suggest that the titanate and acid-exchanged structures were closely related to the pseudolepidocrocite arrangement (\(\text{C}_8\text{Ti}_6\text{O}_{12}\) type). The average structure of NaTi$_2$O$_4$(OH) nanowires consists of noncorrugated Ti$_2$O$_3$(OH) and NaO layers stacked along the [010] direction and an energy minimization was further conducted by theoretical methods. From TEM study, we demonstrated that NaTi$_2$O$_4$(OH) exhibits a modulated structure (q wave vector equals to 1/7 [001]), which originates from the misfit between the NaO and Ti$_2$O$_3$(OH) sublattices. NaTi$_2$O$_4$(OH) structure has a monoclinic distortion (\(\text{P}2/m\) space group) with refined unit cell \(a = 0.37000(6)\) nm, \(b = 1.7816(4)\) nm, \(c = 2.2077(3)\) nm, \(\beta = 92.70(1)\)°. Proton exchange followed by heat treatment convert the sodium hydroxo titanate into highly photoactive TiO$_2$ nanowires. The activity is attributed to the presence of preferentially exposed (001) surface plane which is explained by the topotactic phase transformation of TiO$_2$(B) into anatase.

**1. Introduction**

Because of the potential applications in the fields of energy conversion and environmental protection, the photochemistry of TiO$_2$ is a fast growing area both in terms of research and commercial activity.\(^1,2\) In such applications, the performance of titanium oxide can be optimized with specific nanostructural control over the morphology of the material. A. Selloni et al. reported lately that the dissociation of water spontaneously occurs on (001) and defected (101) surfaces of anatase TiO$_2$ according to the calculated results.\(^3\) This implies that Titanium oxide nanomaterials with preferred orientation and high active site density on the (001) and (101) surfaces could significantly enhance the photocatalytic efficiency. Recently, various titanium oxide nanowires\(^4\) and nanotubes,\(^7,8\) with preferred orientations, have been synthesized from sodium titanate via chimie douce (soft chemistry) routes. Among them, TiO$_2$(B) nanowires, exhibiting excellent photocatalytic efficiency, even better than P25 Degussa and commercial anatase, have been synthesized.\(^9\) In literature, numerous methods have been reported for the synthesis of sodium titanate nanowires with complex structures.\(^4\)\(\cdots\)\(^9\)\(\cdots\)\(^11,15\) Many of them,\(^9\)\(\cdots\)\(^14\) have been converted into titanium dioxide under...
appropriate reaction conditions. However, the exact structures of the titanates remain largely misunderstood. Several standard structural models have been proposed as assumed intermediates, namely layered titanates such as Na₂Ti₂O₅, Na₂Ti₃O₇, or lepidocrocite types. Zhu et al. proposed that the structure of the nanowire is a lamellar arrangement as in the tetratitanate structure, whereas Wu et al. suggested that nanowires are composed of trititanate units. A lepidocrocite structure, with infinite zigzag chains, has also been assigned as nanowire constituent. More understanding of the structure of titanate nanowires may be helpful to comprehend their growth mechanism and to provide tailored synthesis of TiO₂ with enhanced properties. In this work, we report a thorough structural characterization of sodium titanate nanowires prepared by a chimie douce method. Further processing of the nanowires generates TiO₂(B) and anatase TiO₂ nanowires. We found that this anatase nanowires with highly (001)-exposed shows enhanced photocatalytic activity better than Degussa P₂₅ TiO₂, which is the currently reported best commercial photocatalyst. This result strongly supports the theoretical predictions provided by A. Selloni.

2. Experimental Section

Preparation of Na₂Ti₄O₉(OH)₂·H₂O·yCO₂ A. Titanate nanowires were synthesized by adding TiO₂—anatase (1.0 g, Riedel-de-Haën) or amorphous TiO₂(1.0 g, synthesized from TiOCl₂ hydrolysis by excess NH₄OH) to a 10 mol L⁻¹ aqueous solution of NaOH (10 mL). The mixture was transferred to a Teflon-lined autoclave and heated to 423 K for 24–72 h. The material was washed with distilled water then filtered. The steps were repeated until pH becomes lower than 12. After centrifuging and drying at 343 K in air, the material was collected by centrifuging repeatedly until pH is above 6. After being washed with distilled water then filtered, the steps were repeated until pH becomes lower than 12. After centrifuging and drying at 343 K in air, A (1.3 g) was isolated.

Preparation of Na₂Ti₄O₉(OH)₂·H₂O·yCO₂ B. To NaOH(aq) (10 mol L⁻¹, 30 mL) in a 300 mL PTFE round-bottom flask equipped with a coil condenser was added amorphous TiO₂(1.0 g, synthesized from TiOCl₂ hydrolysis by excess NH₄OH). The mixture was refluxed in an air atmosphere at 423 K (oil bath temperature) for 72 h. B (1.3 g) was isolated via the analogous workup procedure mentioned above.

Preparation of HTi₂O₄(OH)·xH₂O·yCO₂ A and HB. The A (1.0 g), product was acid-washed, which involved stirring the sample in 0.1 mol L⁻¹ HNO₃(aq) solution (50 mL) for 2 h. The material was then filtered, washed with distilled water, and collected by centrifuging repeatedly until pH is above 6. After being dried in an oven at 343 K under air, HA (0.8 g) was obtained. HB can be prepared by using the same process.

Preparation of TiO₂(B) HA-1 and HB-1. TiO₂(B) nanowires, HA-1 and HB-1, were prepared by heating the acid-washed titanate nanowires, HA and HB, respectively, at 673 K (rate, 10 K min⁻¹), for 4 h in air.

Preparation of Anatase TiO₂, HA-2 and HB-2. Subsequent thermal treatment of HA and HB at 1073 K (rate, 10 K min⁻¹) for 4 h in air, converts them to HA-2 and HB-2, respectively.

Preparation of Sodium Titanate, Titanic Acid, and TiO₂ Nanotubes. Titanate nanowires were synthesized by adding TiO₂—anatase (1.0 g, Riedel-de-Haën) to a 10 mol L⁻¹ aqueous solution of NaOH (10 mL). The mixture was transferred to a Teflon-lined (50 mL) autoclave and heated to 423 K for 24–72 h. The material was washed with distilled water following a procedure similar to that of the A preparation and then filtered. For preparation of titanic acid and TiO₂ nanotubes, the above isolated powder was treated in a process similar to that of HA and HA-1 preparation, respectively.

Characterization. SEM images were obtained by using a JEOL 6400F with a tungsten cathode field emission gun operating at 10 kV. XRD data were collected on a Siemens D5000 diffractometer with the Cu Kα radiation in a Bragg–Brentano geometry. TEM studies were performed on a Hitachi HF2000 with a field emission electron source at 200 keV. Samples prepared by grinding solid products followed by dispersing them in ethanol were placed on holey or lacy carbon films on copper grids for TEM studies. TEM simulations were performed using a specialized program “Mac Tempas”, which uses parameters of the structures, including cell parameters, space group, atom positions, thickness, and defocalization. EDX studies were performed by using a PGT X-ray analysis probe attached to a JEOL 5800LV SEM with a conventional tungsten electron source operating at 15 keV. The EDX samples were prepared by pressing 100 mg of powder into a pellet with 8 mm diameter and 1–2 mm thickness. The signal was collected from a well-planar surface to limit the differential absorption phenomenon. In addition, a standard sample NaCl is used to carefully calibrate the detector and the decomposition procedure. In that case, the accuracy was estimated lower than 3% for the Na/Ti molar ratio. TGA-MS studies were carried out using a SETARAM TG-DSC 111 equipment fitted with Mass Spectrometer facility (Leybold Inficon) operating at 273–1073 K with a heating rate of 5 K/min, in Ar environment. FT-IR spectra were observed using a Bruker Vertex 70 equipped with a universal attenuated total reflection (ATR) accessory (Golden Gate™ Single Reflection Diamond ATR). FT-Raman studies were performed on a Bruker RFS100 equipped with a 1064 nm infrared laser as the incident light, which allows one to focus a sample area with a diameter less than 1 mm. The specific surface area determined from the Brunauer–Emmett–Teller equation (BET method) was performed by N₂ adsorption at 77 K (under liquid nitrogen) using the adsorption apparatus Micromeritics ASAP 2010 instrument. The density measurements were carried out at room temperature by means of an ACCUPYC 1330 Pycnometer (Micromeritics) with a high accuracy (for a total volume of 1 mL, AV is lower than 1 μL). Before the measurements, all compounds were prepared in an oven at 423 K for one day.

Photocatalytic Study. Photodecomposition of methylene blue was used to test photocatalytic efficiency of the titanium dioxides prepared in this study. Irradiation was provided by a Xe lamp (Osrain 180W). Photocatalytic tests were performed at room temperature. For each test, TiO₂(10 mg) was added, under stirring, to an aqueous solution of methylene blue (25 ppm, 20 mL). A sample (0.5 mL) was taken from the suspension at a designed time interval for analysis. The methylene blue concentration of samples was determined by Hitachi U3010 UV–vis spectrophotometer.

Computational Parameters. Crystal structure of phase A has been relaxed with the Density Functional Theory (DFT) code VASP. In this code, atoms are represented by plane waves in the Plane Augmented Waves (PAW) formalism. We used the Generalized Gradient Approximation (GGA) with the Perdew, Burke, Ernzerhof (PBE) functional for exchange and correlation

The sample suggests that it is more fragmented than fragmentation is more evident in HA-1 but their lengths decrease to 0.1 HA nanowires in post-treatment at 1073 K, the form-factor (length to width microscopic (SEM) image in Figure 1a shows that A mostly retained in HA-2 and HB-2 at 1073 K, HB-1 HB-1 and HB-2 were isolated, correspondingly. Detailed experimental conditions for samples preparation and morphological properties are summarized in Table 1.

3. Results and Discussion

The treatment of TiO$_2$ precursors in NaOH solution was conducted either in autoclave or refluxing conditions, which produced A and B compounds respectively. The HA and HB samples, in which Na$^+$ ions were fully exchanged by protons, were obtained by washing A and B in diluted acidic solutions. Finally, annealing of HA and HB at 673 K led to HA-1 and HB-1, respectively. Upon calcinations of HA and HB at 1073 K, HA-2 and HB-2 were isolated, correspondingly. Detailed experimental conditions for samples preparation and morphological properties are summarized in Table 1.

3.1. Morphology Study by SEM. A scanning electron microscopic (SEM) image in Figure 1a shows that A is composed of nanowires with diameters 50 – 200 nm and lengths 0.5 – 5 µm. Apparently, the morphology of A is mostly retained in HA (Figure 1b). Close examination of the sample suggests that it is more fragmented than A. The fragmentation is more evident in HA-1 (Figure 1c). After post-treatment at 1073 K, the form-factor (length to width aspect ratio) of HA-2 further decreased. In Figure 1d, the nanowires in HA-2 show almost the same diameters as in HA but their lengths decrease to 0.1 – 1 µm. HA-2 will be denoted nanorods hereafter. Sample B, obtained in refluxing conditions at 423 K, is composed of small nanowires with diameters 10–40 nm and lengths 0.05–1 µm as shown in Figure 1e. Further post-treatments of B led to the formation of HB, HB-1, and HB-2 (Figure 1f). They exhibit a morphology variation trend analogous to that observed for the samples treated from A. In both cases, increasing reaction duration increased the lengths of the nanowires only. The diameters essentially remained the same. This observation is in agreement with previous reports.6,9

3.2. Composition Determination. Using an energy-dispersive X-ray (EDX) analyzer attached to the SEM, we found that A and B contained a significant amount of Na, with Na/Ti ≈ 0.52, whereas HA and HB were essentially Na free. The IR spectra of A, B, HA, and HB showed absorptions from vibrations of O–H and Ti–O bonds (Figure S1 in the Supporting Information). In addition, the spectra of A and B displayed C–O bands, which are assigned to the presence of carbonate groups, νC–O$_I$ and νC–O$_{II}$ at 1380 and 1680 cm$^{-1}$, respectively. For HA and HB, intensities of the carbonate bands were much weaker, whereas the OH bands remained unchanged. The νC–O$_I$ vibration attributed to undentate carbonates is no more observed. This probably was the result of the acidification of A and B. Thermogravimetric analysis/mass spectrometry (TGA/MS) was undertaken to estimate the quantity of the bonded hydroxyl and carbonate groups, and the adsorbed water molecules in the samples (Figure S2 in the Supporting Information). For all samples, two water departures below

<table>
<thead>
<tr>
<th>sample</th>
<th>precursor</th>
<th>preparation</th>
<th>morphology</th>
<th>dimension (width/length (nm))</th>
<th>phase</th>
<th>specific surface (m$^2$ g$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>RDH TiO$_2$ d: 100</td>
<td>10 M NaOH$_{aq}$, 453 K, 48 h in autoclave</td>
<td>fibrous</td>
<td>50–200/500–5000</td>
<td>lepidocrocite: &quot;Cs$_2$Ti$<em>6$O$</em>{12}$&quot;-type</td>
<td>27</td>
</tr>
<tr>
<td>HA</td>
<td>A</td>
<td>0.1 M HCl$_{aq}$, 298 K, 2 h</td>
<td>fibrous</td>
<td>50–200/500–5000</td>
<td>lepidocrocite: &quot;Cs$_2$Ti$<em>6$O$</em>{12}$&quot;-type</td>
<td>52</td>
</tr>
<tr>
<td>HA-1</td>
<td>HA</td>
<td>673 K, 4 h</td>
<td>fibrous</td>
<td>50–200/100–5000</td>
<td>TiO$_2$(B)</td>
<td>41</td>
</tr>
<tr>
<td>HA-2</td>
<td>HA</td>
<td>1073 K, 4 h</td>
<td>fibrous</td>
<td>50–200/100–2000</td>
<td>lepidocrocite: &quot;Cs$_2$Ti$<em>6$O$</em>{12}$&quot;-type</td>
<td>35</td>
</tr>
<tr>
<td>B</td>
<td>amorphous TiO$_2$</td>
<td>reflux in 10 M NaOH$_{aq}$, 48 h</td>
<td>fibrous</td>
<td>10–40/50–1000</td>
<td>lepidocrocite: &quot;Cs$_2$Ti$<em>6$O$</em>{12}$&quot;-type</td>
<td>105</td>
</tr>
<tr>
<td>HB</td>
<td>B</td>
<td>0.1 M HCl$_{aq}$, 298 K, 2 h</td>
<td>fibrous</td>
<td>10–40/50–1000</td>
<td>TiO$_2$(B)</td>
<td>71</td>
</tr>
<tr>
<td>HB-1</td>
<td>HB</td>
<td>673 K, 4 h</td>
<td>fibrous</td>
<td>10–40/50–1000</td>
<td>TiO$_2$(B)</td>
<td>25</td>
</tr>
<tr>
<td>HB-2</td>
<td>HB</td>
<td>1073 K, 4 h</td>
<td>fibrous</td>
<td>10–40/50–1000</td>
<td>anatase</td>
<td>262</td>
</tr>
<tr>
<td>HB-nanotube</td>
<td>RDH TiO$_2$ d: 100</td>
<td>10 M NaOH$_{aq}$, 423 K, 48 h in autoclave</td>
<td>tubular</td>
<td>5–10/50–300</td>
<td>sodium titanate</td>
<td>284</td>
</tr>
<tr>
<td>TiO$_2$ nanotube</td>
<td>HB-nanotube</td>
<td>0.1 M HCl$_{aq}$, 298 K, 2 h</td>
<td>tubular</td>
<td>5–10/50–300</td>
<td>less-ordered titanic acid</td>
<td>323</td>
</tr>
<tr>
<td>Degussa P25</td>
<td>673 K, 4 h</td>
<td>fibrous</td>
<td>5–10/10–100</td>
<td>anatase + TiO$_2$(B)</td>
<td>50</td>
<td></td>
</tr>
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</table>

"synthesized from hydrolysis of TiOCl$_2$ in excess NH$_4$OH.

Figure 1. SEM images of (a) A, (b) HA, (c) HA-1, (d) HA-2, (e) B, and (f) HB-2.
structures originates from the relative stacking of the ribbons (along the [010] direction), which is built up from antiphased layers along the a axis for “Cs2TiO1.6” type, instead of phased layers in lepidocrocite (Figure S3 in the Supporting Information).

Referring to the rock-salt structure, “Cs2TiO1.2” can be described as a periodic stacking of three layered sublattices, i.e., [□□□], [□O], and TiO, along [010] in which the Cs atoms occupied the oxygen vacancy sites (noted □). In terms of NaCl-type, the deduced composition is written [Ti2O3][O2□]. Although possible structures of chimie douce titanate nanowires have been proposed to possess (C2/m or P2/m) monoclinic alkali titanates Na2Ti₅O₁₀ or Na2TiO₇ structures, the XRD results did not match well with the known data. The agreement with the Raman data is also not in favor of the tetra- or trititanate. The peak observed at 910 cm⁻¹ is indeed assigned to a dangling Ti–O sticking out into the interlayer space, in a stepped layer structure. Here (Figure S4 in the Supporting Information), this peak is hardly marked and this observation implies that the structures are similar to a noncorrugated layer or a tunnel structured titanates. All the evidence reveals that A does not have a typical stepped layer structure. Therefore, we propose that a structural model for A could derive from a noncorrugated layered structure closely related to “Cs2TiO1.6.” Detailed structural description for HA will be discussed in section 3.3.4. The XRD patterns of HA-1 and HA-2 in Figure 2 suggest that the allotropic variety is TiO₂(B) and anatase phases, respectively. Additional structural features of both HA-1 and HA-2 will be described in 3.3.4. too. The patterns of B, HB, HB-1, and HB-2 (see Figure S5 in the Supporting Information) exhibit almost the same peak positions and relative intensities, but a broadening in peak widths was detected, as those of the corresponding samples developed from A.

3.3.2. TEM Studies of NaTi2O4(OH). Figure 3a shows a low magnification transmission electron microscopic (TEM) image of A, showing bundles of nanowires with apparent diameters of tens of µm and lengths of several µm. From a separated nanowire in Figure 3a, a high-resolution (HR) image in Figure 3b is obtained. Figure 3c is the filtered image of Figure 3b. The white rectangular area in Figure 3c represents the centered unit cell observed along [010]. In Figure 3b, the distance between the repeating fringe units is 1.1 nm, which significantly differs from stepped-layer unit evidenced in tetra-titanate (1.2 nm) or trititanate (0.9 nm). The selected area electron diffraction (SAED) of A along [010] (Figure 3d) gives evidence for two main characteristic distances 0.37 and 0.315 nm with the following reflection conditions (h00): h = 2n, (00 l); l = 2n, (h0l); h + l = 2n. These observations are not in agreement with the space group extensions of the stepped layer structures such as Na₂Ti₅O₁₀ (C2/m, reflection condition h + k = 2n) or Na₂TiO₇ (P2/m, no special reflection condition) but are similar to those observed in the orthorhombic bronze, Cs₂Ti₅O₁₂ (Immm), which has a noncorrugated layer structure. The (010) reciprocal plane analysis reveals a slightly distorted cell with an angle equal to β* ≈ 87°, which suggests a monoclinic distortion with an I mode. Besides this, a reinforcement of the intensity every 7 spots can be seen on the ED pattern, along the [001]* direction. The existence of six satellites peaks along [001]* suggests a structural modulation in which...
the q wave vector equals $1/n[001]^\circ$. The lattice parameter of the superstructure is then $c = nc'$, i.e., $7 \times 3.15 \approx 2.2$ nm, where $c'$ is close to the $c$ parameter of “Cs$_2$Ti$_6$O$_{12}$” unit cell, 0.2981(2) nm. TEM and XRD analysis show that $A$ crystallizes in a monoclinic system, $I\overline{2}/m$ space group, with the following refined cell parameters $a = 0.37000(6)$ nm, $b = 1.7816(4)$ nm, $c = 2.2077(3)$ nm, $\beta = 92.70(1)^\circ$. The refinement was conducted using Jana 2000. The peak position and intensity of the experimental and the refined XRD patterns with reliability factors are summarized in the Supporting Information (Figure S6 and Table S1).

To support this analysis, we modeled the structure of NaTi$_2$O$_4$(OH) to be analogous to that of “Cs$_2$Ti$_6$O$_{12}$”, preserving the TiO framework. Cell parameters and atomic positions were fully relaxed starting from the XRD refinement results, with a force convergence criterion of $5 \times 10^{-3}$ eV Å$^{-1}$. The relaxed structure is in remarkable agreement with the averaged pseudolepidocrocite structure since we obtained the following cell parameters $a = 0.364$ nm (1.58%), $b = 1.729$ nm (2.97%), $c = 0.302$ nm (3.96%), $\alpha = 91.2^\circ$, $\beta = 89.9^\circ$ and $\gamma = 88.2^\circ$ (relative errors with respect to the XRD parameters are reported in parenthesis). Relative error for cell parameters remains below 4% and it is worth noting that the highest discrepancy between experimental and calculated cell parameters is for $c$ parameter that is in the direction of the structural modulation. The position of atoms in the proposed structure is only preliminary (see the Supporting Information, Table S3) and further calculation of energy minimization and XRD refinement will be necessary to take into account the modulation.

3.3.3. Modulated Layer Structure of NaTi$_2$O$_4$(OH). Assuming that the misfit between NaO and TiO sublattice distances causes the observation of six satellite peaks on the ED pattern along [001]$^\circ$, we can propose the structural model shown in Figure 4. This assumption is corroborated by the fact that the sodium hydroxide (NaOH noted hereafter SH) structure, in beta form, also contains rock-salt subunits with a periodicity $a_{SH} = 0.3345(3)$ nm and $b_{SH} = 0.3445(3)$ nm in the (110) plane. Compared to the TiO lattice $a = 0.3821(3)$ nm, $c = 0.2981(2)$ nm in “Cs$_2$Ti$_6$O$_{12}$”, and without (28) Petricek, V. The Crystallographic Computing System JANA2000; Institute of Physics: Prague, Czech Republic, 2000.
any accommodation of this misfit, the Na–O lattice is in tension in the [100]_{\text{Cs}_2\text{Ti}_6\text{O}_{12}} direction whereas it is compressed in the [001]_{\text{Cs}_2\text{Ti}_6\text{O}_{12}} direction. One possibility to accommodate the misfit is to displace the Na and O atoms along the a axis in the (a, c) plane (Figure 4a–c). This misfit could be reduced if a sinusoidal distortion of the Na and O atoms is applied into the (101) plane to accommodate the Na–O distances at 0.337–0.344 nm values. Along the c-axis, the sequence repeats every seven regular Ti–O sublattices which becomes in coincidence with the distorted Na–O sublattice. Such a model is coherent with the seventh order superstructure found in the [010]∗ zone axis in Figure 3d. These results give evidence that the NaTi2O4(OH) nanowires exhibit a commensurate modulated structure along the [001] direction. Figure 4d shows the proposed structural model along the [212] direction. This representation gives an idea of the crystalline directions that are preferentially developed during the nanowire growth.

For further comparison of the assumed structure and the TEM observations, Mac Tempas simulation of ED (see Figure S10 in the Supporting Information) and HRTEM patterns were performed on an idealized structural model of a monoclinic system with R2/m space group. As shown in Figure S10b, the peaks intensity and position of the simulated ED pattern, viewed along [010]∗, qualitatively agree with the experimental result. In Figure 3c, the HRTEM image simulated from the idealized model (focal = 90 nm; thickness 34 nm) viewed along [010]∗ has been found to superimpose well on the observed image. The modeled structure is coherent with the other experimental results: EDX chemical composition analysis, theoretical and experimental densities (d_{calc} = 3.20; d_{obs} = 3.17(2)) and weight loss measured by thermogravimetric analysis.

3.3.4. Characterization of HTi2O4(OH), TiO2(B), and Anatase TiO2. As shown in Figure 2, the XRD pattern of HA is slightly different from that of A. In particular, only the most intense peaks are still present in HA, the product acidified from A, whereas the peak widths are broader than those of A. Referring to the previous indexations of A and assuming a layered structure for the fully proton exchanged acidified from TiO2(B), the two main peaks of A are identified as (002) and (020) (interlayer spacing). In a first approximation, the refined parameters $a = 0.37405(2)$ nm, $b = 1.67444(4)$ nm, $c = 2.2143(2)$ nm, $\beta = 92.70(1)^\circ$ indicate a significant contraction of the interlayer spacing while the other parameters are slightly changed. Such behavior is in agreement with that usually observed for acid exchanged titanates. The peak position and intensity of the experimental and the refined XRD patterns with reliability factors are shown in Table S4 of the Supporting Information. For comparison, all the intensity values were normalized to the (112) peak intensity of nonoriented anatase, the nature of adsorbed species (HCOO–, CO2–) strongly impact the final form of materials. Typically, (101) and (001) surface planes are found in P25 powder materials, together with some (100)/(010). The anatase nanowires, obtained by “chimie douce”, mainly exhibit (001) surface plane and a growing axis along the [010] direction is observed in TEM images. To quantify the preferential orientations, we performed an analysis of the X-ray diffraction peaks (Figure S8 in the Supporting Information). The XRD peak intensities of HA-1, HB-1, HB-2, P25, TiO2-tube are summarized in Table S4 of the Supporting Information with calculated intensities of randomly oriented anatase (JCPDS 89–4921, calculated intensities). The intensity values were normalized to the (112) peak intensity of nonoriented anatase, denoted reference or ref, in the following. The preferential orientation can be estimated through the ratio between the XRD peak intensities of the samples to the same peak intensities of the reference, $I_{\text{sample}}/I_{\text{ref}}$ (see Table 2). The $I_{(101)}/I_{(101)\text{ref}}, I_{(004)}/I_{(004)\text{ref}}$ and $I_{(200)}/I_{(200)\text{ref}}$ ratios for HA-1 (autoclave) and HB-2 (reflux) are stronger than that of P25. Therefore, the anatase nanowires, converted from the titanate nanowires, expose more (101), (001), and (000) planes than the other samples. In comparison to HA-1, HB-2 shows a higher (004) peak intensity, whereas their respective (101) and (200) intensities are almost the same. The preferential orientation in the (001) plane and consequently the increased

![Figure 5](image)

impact the photocatalytic properties (see below). The shape and size of the nanocrystallites vary with the preparation techniques and procedures. Factors such as ionic strength, nature of the ligand (aquo, oxo, or hydroxo) at the surface of the reactant (amorphous or crystalline precursor), thermal treatment (autoclave (A), reflux (B)), and in addition, the nature of adsorbed species (HCOO–, CO2–) strongly impact the final form of materials. Typically, (101) and (001) surface planes are found in P25 powder materials, together with some (100)/(010). The anatase nanowires, obtained by “chimie douce”, mainly exhibit (001) surface plane and a growing axis along the [010] direction is observed in TEM images. To quantify the preferential orientations, we performed an analysis of the X-ray diffraction peaks (Figure S8 in the Supporting Information). The XRD peak intensities of HA-1, HB-1, HB-2, P25, TiO2-tube are summarized in Table S4 of the Supporting Information with calculated intensities of randomly oriented anatase (JCPDS 89–4921, calculated intensities). For comparison, all the intensity values were normalized to the (112) peak intensity of nonoriented anatase, denoted reference or ref, in the following. The preferential orientation can be estimated through the ratio between the XRD peak intensities of the samples to the same peak intensities of the reference, $I_{\text{sample}}/I_{\text{ref}}$ (see Table 2). The $I_{(101)}/I_{(101)\text{ref}}, I_{(004)}/I_{(004)\text{ref}}$ and $I_{(200)}/I_{(200)\text{ref}}$ ratios for HA-1 (autoclave) and HB-2 (reflux) are stronger than that of P25. Therefore, the anatase nanowires, converted from the titanate nanowires, expose more (101), (001), and (000) planes than the other samples. In comparison to HA-1, HB-2 shows a higher (004) peak intensity, whereas their respective (101) and (200) intensities are almost the same. The preferential orientation in the (001) plane and consequently the increased

the stability of TiO2(B) could be assigned to linked carbonate contains also some amount of carbonates, suggesting that the processed material crystallizes as anatase mixed with traces of TiO2(B). The intensities of (101) and (004) peaks show that anatase is more present compared to the non-oriented anatase phase. The (200) peak is not observed, indicating a lower percentage of [100] orientation along the nanotubes. The proton-exchanged TiO2 nanotubes after proton exchange maintained the tubular morphology but the wall structure became less ordered. The highly strained scroll structure of the nanotube before and after the ion-exchange process appears through a (020) peak broadening in the XRD pattern. The proton-exchanged TiO2 nanotubes were further annealed at 673 K. As shown in the XRD pattern (Figure S9d in the Supporting Information), the processed material crystallizes as anatase mixed with traces of TiO2(B). The intensities of (101) and (004) peaks are 13% and 6% lower while that of (200) is 13% higher than the non-oriented anatase. This suggests that the anatase TiO2 processed from the nanotubes may have numerous stacking faults in the (020) planes and a slightly preferential orientation along (100).

### 3.3.5. TiO2(B) to Anatase Phase Transformation

In comparison to the literature, the transformation of TiO2(B) to anatase, which generally occurs at T = 823 K as for the HB-1 to HB-2, takes place at 1073 K for the HA-1 to HA-2. The indicated temperatures refer to the m = 44 peak observed by MS (Figures S2c and S2d in the Supporting Information) and attributed to CO2 departure. A major difference between our work and previous ones mainly lies in the size of the fibers. The classical synthesis leads to micrometric crystals: 0.25–0.7 μm wide; 1–5 μm long, whereas small or large nanowires are formed by the low temperature process. In addition, the transformation is associated with the release of carbonate groups (probably bidentate), as observed by TGA/MS experiments (Figures S2c and S2d in the Supporting Information). Such an observation may indicate that these adsorbed species also play a critical role in the solid state crystal transformation. Therefore, the same analysis (TGA/MS) was performed on micrometric fibers of TiO2(B). It clearly appears that the latter contains also some amount of carbonates, suggesting that the stability of TiO2(B) could be assigned to linked carbonate groups on the crystallite surface which may stabilize the TiO2(B) surface and create an additional barrier which hampers the phase transformation process toward anatase. However, the higher thermal stability of the present HA-1 TiO2(B) can be attributed both to the presence of carbonate and to the size and morphology of the crystallites. This conclusion is further supported by the fact that the quantity of carbonate molecules decreases as the size of the crystallites increases. The transition from TiO2(B) to anatase occurs at a lower temperature for HB-1 small nanowires with largest surface area (71 m2 g−1), whereas the higher one is observed for HA-1 with large nanowires and smaller surface area value (41 m2 g−1).

### 3.4. Structural Reconstruction Mechanism

According to the observations discussed above, we propose a possible mechanism in Scheme 1 to rationalize the formation pathways of the titanates and the TiO2 compounds. In hot concentrated base solution, the starting TiO2 particles are dissociated into building blocks composed of linked titinate units. They then recombine into pseudolepidocrocite sodium hydroxo titanate nanowires with the long axis oriented along the a direction. This particular arrangement is strongly depending on the Na content, because for lower Na/Ti ratio, the reported Na titanate could not be stabilized. The Na ions in the titinate nanowires can be totally proton exchanged in diluted acid solution and the pseudolepidocrocite structure is retained. After annealing at 673 K, dehydration and condensation of the layered titanic acid convert it into TiO2(B) nanowires with growth direction along [010] axis. As shown in Scheme 1, the condensation step might proceed via connecting the same colored units after simple shifting of the different colored units. By another simple TiO6 octahedron migrations shown in the final step in Scheme 1, anatase can be formed from the TiO2(B) intermediate. The detailed mechanism for the latter phase transformation has been widely studied.30 The as-prepared anatase crystallizes as nanowires with axis-direction along [010] and exposed (001), (100), (101) planes. The octahedron units migrating in the last step are emphasized by marking them in bold lines. The (001) and defective (101) surfaces were predicted to be photoactive.3

Varying the reaction conditions (Table 1) influences the product morphology in many ways. For example, employing amorphous TiO2 instead of anatase TiO2, as starting material in the autoclaved process generated thinner diameter sodium titanate nanowires. On the other hand, using anatase TiO2 in the refluxing process produced sodium titanate nanotubes. The dissociation/dissolution reactions of precursor would be probably more efficient at higher temperature as in autoclave process than that in reflux conditions which operate at lower temperature. Beside this, the sizes of the building blocks, fragmented from the precursor solids, correlate strongly to the size and morphology of the final products. In another case, B, which shows analogous phase conversion behavior as A does, displays smaller diameters. The reason for this might be the combination of several reaction conditions, such as smaller sized initiator, lower reaction temperature, and lower reaction pressure. From the topotactic condensation of titanate layers proposed in Scheme 1, the thinner nanowires HB-2 can form, exposing more (001) plane but less of (100) (011). 

Table 2. Summary of XRD Relative Intensities, Iref/Iobs, Together with Photocatalytic Property of the Samples

| Sample | HA-1 | HA-2 | HB-1 | HB-2 | TiO2 nanotube | HA-1 | HA-2 | HB-1 | HB-2 | P25 |
|--------|------|------|------|------|-------------|------|------|------|------|-----|------|
| I(101)/I(100) | 1.10 | 1.40 | 1.36 | 0.50 | 0.60 | 1.06 | 1.72 | 1.42 | 2.00 | 0.83 | 0.94 | 1.13 | 1.00 |
| I(004)/I(100) | 1.36 | 0.50 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 | 0.93 | 1.23 | 1.13 | 1.50 |

* Rations of the XRD intensities of chosen anatase planes normalized to non-oriented anatase. JCPhysD 89–4921. Initial reaction rate estimated in first step of the photodecomposition.
estimation corresponds well with the structural observations of HA-2 and HB-2.

3.5. Photocatalysis Study. Photocatalytic activities of TiO₂(B) nanowires HA-1, HB-1, anatase nanowires HA-2, HB-2, TiO₂ from the nanotubes (TiO₂-nanotube) and Degussa P25 nanoparticles were examined in the photodecomposition of an aqueous solution of 25 ppm methylene blue. The kinetic curves are shown in Figure 6 and the initial reaction rates \( r_0 \), in the first step of photodecomposition, are summarized in Table 2. Interestingly, the efficiency of the methylene blue photodecomposition, deduced from \( r_0 \) values, decreases in the following order: HB-2 > P25 > TiO₂-nanotube > HB-1 > HA-2 > HA-1. HB-2 displays an efficiency 25 and 75% higher than that of P25 and HA-2, respectively. According to the specific surface in Table 1, the surface area of HB-2 (25 m² g⁻¹) is half that of P25 (50 m² g⁻¹) and twice that of HA-2 (13 m² g⁻¹). From the previously mentioned observations, in section 3.3.4., HB-2 and HA-2 expose more (101), (001), and (100) planes than do other TiO₂ anatase. Although HA-2 and HB-2 crystallize as anatase nanowires, HA-2, because of its thicker crystallite-size and lower surface area, exhibits less exposed (001) surface planes than HB-2 does, as deduced from the XRD study. Therefore, the higher activities of HB-2 can be attributed to more exposed (001) than (100) and (101) surface planes, whereas the good activity of P25 might originate from exposed (001) and (100) planes and its higher surface area. This observation is in agreement with the calculated results, i.e., the (001) surface of anatase is more active than (101).³ Further investigations in that field are needed to produce larger anatase nanowires with (001) preferential orientation.

Even though the TiO₂-nanotube prepared in this study exhibits the highest specific surface area, 323 m² g⁻¹, its rather average photocatalytic activity, with \( r_0 \) being 1.0 ppm min⁻¹, is lower than those of HB-2 and P25. The nanotube shows less exposed (001) and (101) surface planes, in addition the inactive (100) surface is more extended than in nonoriented anatase. All the observations mentioned above give a strong experimental evidence and successfully validate the calculated results regarding anatase surface activity³ and explain why HB-2 has excellent photocatalytic activity. In future works, to reduce the nanowire thickness will be the key point to improve photocatalytic activity of the products. Although the photocatalytic activity of TiO₂(B) nanowires has been reported to be higher than that of P25 Degussa,³ in this study, HA-1 and HB-1 did not exhibit better efficiency than P25 did.
4. Conclusions

In summary, the structure of NaTi$_2$O$_4$(OH) nanowires and derivative products prepared by the alkali method (reflux and autogenous pressure) were studied in details using XRD, ED and HRTEM. Based on density measurements, crystallographic, EDS and TG/MS results, we confirm that the titanate nanowires consist of pseudolepidocrocite arrangement ("Cs$_2$Ti$_6$O$_{12}$" type) rather than Na$_2$Ti$_3$O$_7$, as previously reported.$^{31,32}$ We demonstrated that NaTi$_2$O$_4$(OH) exhibits a modulated structure ($q$ wave vector equals 1/7 [001]*), which originates from a misfit between the NaO and Ti$_2$O$_3$(OH) sublattices. NaTi$_2$O$_4$(OH) structure presents a monoclinic distortion ($I\overline{2}/m$ space group), with refined parameters $a = 0.37000(6)$ nm, $b = 1.7816(4)$ nm, $c = 2.2077(3)$ nm, and $\beta = 92.70(1)^\circ$. This work contributes to clarify the current controversy on the structures of Na-titanates nanowires. Proton exchange of NaTi$_2$O$_4$(OH) followed by heat treatment converts it into titanium dioxide nanowires with TiO$_2$(B) and then anatase structures. The proposed mechanism of anatase nanostructure formation elucidates the relationship between 2D and 3D nanowire structures. The long axis of the anatase nanowires oriented along the [010] direction and the (001) preferential orientation was explained by a topotactic phase transformation of TiO$_2$(B) into anatase. The length and thickness of nanowires along (100) can be controlled by adjusting the reaction temperature. The higher photoactivity of anatase TiO$_2$ nanowires, produced by reflux process, originates from the (001) preferential orientation, despite its lower surface area, as compared to P25. In future works, to reduce the nanowire thickness will be the key point to improve photocatalytic activity of the products.

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Supporting Information Available: TGA/MS, FT-IR, Raman, and XRD of A and H-A (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.