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Introduction

Heterogeneous solid catalysts have been widely used in various chemical reactions in the chemical industry, exhaust gas puri fication, and environmentally friendly reactions. They are more suitable for reactions at high temperatures than homogeneous catalysts, and are removed easily from the reactor. In particular, solid acid catalysts are used in many important processes related to petroleum refining and petrochemical production. Consequently, numerous studies have specifically examined solid acid catalysts. On the other hand, fewer efforts have been devoted to heterogeneous base catalysts.¹ Because general base sites on a solid base catalyst are readily poisoned by moisture and carbon dioxide in the atmosphere, 2 the development of a promotion strategy for solid base catalysts is more difficult than for solid acid catalysts.

Layered double hydroxide, generally called LDH, is a wellknown layered clay mineral that is known to act as a unique solid base catalyst. Actually, LDH comprises brucite-like positively charged two-dimensional sheets denoted as $[M_{1-x}^{2+x}M_x^{3+(OH)_2}]^{x+x}$ and interlayer parts denoted as $A_{x/n}^n$ ⁿ⁻· mH_2O , where A^{n-} corresponds to interlayer anions such as carbonate and hydroxide. The

Effect of $SiO₂$ amount on heterogeneous base catalysis of SiO₂@Mg-Al layered double hydroxide[†]

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The effects of $SiO₂$ amount on the base catalysis of highly active finely crystallized Mg–Al type layered double hydroxides prepared by the co-precipitation method with coexistence of $SiO₂$ spheres, denoted as $SiO₂GLDHs$, were investigated. With the Si/(Mg + Al) atomic ratios of 0–0.50, the highest activity for the Knoevenagel condensation was observed in the case of $Si/(Mg + Al) = 0.17$, as the reaction rate of 171.1 mmol g(cat) $^{-1}$ h $^{-1}$. The base activity increased concomitantly with decreasing LDH crystallite size up to Si/(Mg + Al) atomic ratio of 0.17. However, above the Si/(Mg + Al) atomic ratio of 0.17, the reaction rate and TOF_{base} were decreased although the total base amount was increased. Results of TEM-EDS and ²⁹Si CP-MAS NMR suggest that the co-existing $SiO₂$ causes advantages for dispersion and reduction of the LDH crystallite to improve the base catalysis of $SiO₂@Mg–Al$ LDH, whereas the excess $SiO₂$ species unfortunately poisons the highly active sites on the finely crystallized LDH crystals above a $Si/(Mg + Al)$ atomic ratio of 0.17. According to these results, we inferred that the amount of spherical $SiO₂$ seeds in the co-precipitation method is an important factor to increase the base catalysis of SiO₂@LDHs; *i.e.* the control of Si/(Mg + Al) atomic ratio is necessary to avoid the poisoning of highly active base sites on the LDH crystal. PAPER
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positively charged sheets and the interlayers are alternately laminated to compensate for the charge between sheets.³⁻⁵ Base sites on LDH are mainly regarded as identical Brønsted basic OH⁻ and $HCO₃⁻$ anions, which are adsorbed onto the LDH surface. These unique basic sites can act even in an air atmosphere and can exhibit basic characteristics without pretreatment.⁴ As a solid base catalyst, LDH is well-known to catalyze various organic transformations such as aldol condensation,⁶⁻⁹ Knoevenagel condensation,¹⁰⁻¹³ epoxidation,¹⁴⁻¹⁶ and transesterification.¹⁷⁻¹⁹ Recent studies have indicated that LDHs can promote advanced environmentally friendly reactions such as biomass-derived saccharide conversion^{13,20–25} and photocatalytic conversion of $CO₂$ in an aqueous solution.²⁶–²⁸ Therefore, the development of highly active LDH catalysts is eagerly sought.

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The main base sites on the LDH surface are generally regarded as adsorbed anions located at the corner and edge of a crystal.²⁹ However, the anions in the interlayer space cannot participate in the chemical reactions because of the high charge density of the LDH layers and the high contents of anionic species and water molecules, resulting in strong interlayer electrostatic interactions between the sheets.4,30 Therefore, the delamination of LDH nanosheets³⁰⁻³⁶ and the fine crystallization of LDH on appropriate carriers³⁷⁻⁴² have been conducted to increase the number of exposed active base sites. As-prepared LDH materials also have been evaluated carefully to assess their characteristics and utility as photocatalysts and electrocatalysts,43,44 high active base catalysts for Knoevenagel condensation⁴⁵ and epoxidation,⁴⁶ magnetic separation of proteins,³⁷ pseudocapacitance,³⁸ flame retardancy of epoxy

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resins,⁴⁰ and as adjuvants.⁴¹ Nevertheless, no report describes a study of improvement of base catalysis of LDH itself by fine crystallization followed by in situ growth method of $SiO_2@LDH$ nanoparticles.

An earlier study revealed a co-precipitation method for preparation of small-crystallized LDH catalysts on $SiO₂$ nanospheres and explored the superior base catalyses for the Knoevenagel condensation of benzaldehyde compared with conventional LDHs.⁴⁷ This method is applicable for the preparation of $SiO_2@LDH$ nanoparticles with various compositions and element combinations: *i.e.* SiO₂@M²⁺-M³⁺LDH (M²⁺: Mg²⁺) or Ni^{2+} , M^{3+} : Al^{3+} or Ga^{3+} , and M^{2+}/M^{3+} : 1 or 3). Various characterizations of $SiO_2@LDH$ nanoparticles using XRD, TEM-EDS, and 29Si CP-MAS NMR techniques revealed that the co-existence of small $SiO₂$ sphere (ca. 40 nm diameter) surface generated the starting points of LDH growth via Si–O–M covalent bond formation, leading to the formation of fine-crystallized LDH and enhancement of base catalysis for the Knoevenagel condensation of benzaldehyde with ethyl cyanoacetate. However, the roles of Si-O-M covalent bonds in the finecrystallization of LDH and base catalysis have not been explored well. Therefore, in this paper, we investigated the base properties and structural parameters of as-prepared $SiO_2@Mg-$ Al LDH materials with $Mg^{2+}/Al^{3+} = 3$, and discussed the base catalysis with different Si to $(Mg + Al)$ ratios to reveal the mechanism of our strategy. Paper

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Experimental

Materials and synthesis of catalysts

Tetraethyl orthosilicate (TEOS), triethanolamine (TEA), and benzaldehyde were purchased from Sigma-Aldrich Corp. Sodium carbonate ($Na₂CO₃$), sodium hydroxide (NaOH) and toluene were supplied by Kanto Chemical Co. Inc. Cetyltrimethylammonium bromide (CTAB), magnesium nitrate hexahydrate $(Mg(NO₃)₂·6H₂O)$, aluminum nitrate enneahydrate $(AI(NO₃)₃·9H₂O)$, benzaldehyde and benzoic acid were obtained from Wako Pure Chemical Industries Ltd. Benzaldehyde was purified by distillation under 0.4 Pa pressure. Ethyl cyanoacetate was purchased from Tokyo Chemical Industry Co. Ltd. and was used without further purification.

Spherical $SiO₂$ (40 nm) was prepared according to descriptions in earlier reports.41,46 First, 96 mmol of TEA and 2 mL of TEOS were combined in a 200 mL eggplant flask. The two-phase mixture was heated in an oil bath at 363 K for 20 min without

stirring. When the mixture was removed from the oil bath, 26.0 mL of an aqueous solution (2.8 wt%) of CTAB pre-heated at 333 K was added immediately as a structure-directing agent in a condensation process. Then, it was stirred continuously for 24 h at room temperature. Thereafter, the resulting mixture was added to 50 mL of ethanol to obtain colloidal aqueous suspension. The obtained precipitate was centrifuged for 5 min at 4000 rpm. After decantation, the sediment was re-dispersed through vigorous stirring in 50 mL of an ethanolic solution of ammonium nitrate (20 g L^{-1}), and then refluxed for 1 h. This procedure was repeated three times. The same operation was performed with a solution of concentrated hydrochloric acid in ethanol (5 $g L^{-1}$) to replace the ammonium ions. The final sediment was washed with ethanol, and then dried in vacuo. The obtained spherical $SiO₂$ powder was calcined at 823 K under 1 L min⁻¹ of air flow for 6 h.

The $SiO_2@(Z)$ LDH catalysts (*Z*: desired $Si/(Mg + Al)$ atomic ratio) were prepared via an in situ co-precipitation method according to a previous report.⁴² Spherical SiO₂ (40 nm) was dispersed in 20 mL of water using ultrasound treatment. After 30 min, 0.96 mmol of $Na₂CO₃$ was added to the solution. Then, after an additional 5 min of sonication was conducted, 19.2 mL of metal nitrate aqueous solution $([Mg] + [Al] = 0.075 M)$ was slowly dropped into the spherical $SiO₂$ dispersed solution, followed by stirring at room temperature. The pH was maintained at 10.0 by an aqueous NaOH solution (1 M) during titration. The obtained suspension was stirred for an additional 1 h. After the resulting paste was filtered, it was washed with 1 L of water and ethanol. Then, it was dried at 383 K overnight. The $\text{Si}/(\text{Mg}^{2+} +$ Al^{3+}) atomic ratios were varied from 0 to 0.50 whereas the Mg/Al atomic ratio was adjusted to 3.

Reaction

Knoevenagel condensation of benzaldehyde with ethyl cyanoacetate was proceeded in a 20 mL Schlenk tube under an N_2 flow (30 mL min^{-1}) . The reaction was performed using 1.0 mmol of benzaldehyde, 1.2 mmol of ethyl cyanoacetate, 10 mg of catalysts and 3 mL of toluene at 313 K. The obtained products were analyzed using a GC-FID (GC-2014, Shimadzu Corp.) equipped with a polar column (DB-FFAP, Agilent Technologies Inc.).

Characterizations

X-ray diffraction patterns (XRD) were collected using a SmartLab (Rigaku Corp.) with a Cu Ka X-ray source (40 kV, 30 mA). The LDH (003) and (110) crystallite sizes were calculated using

Fig. 1 Activities for Knoevenagel condensation of benzaldehyde with ethyl cyanoacetate over as-prepared SiO₂@LDHs with various Si/(Mg + Al) atomic ratio; (A) time-based reaction progression on benzaldehyde conversion and (B) reaction rate. Reaction conditions: benzaldehyde (1.0 mmol), ethyl cyanoacetate (1.2 mmol), catalyst (10 mg), toluene (3 mL), 313 K, N₂ flow (30 mL min⁻¹). Number in (A) denotes Si/(Mg + Al) atomic ratio.

the Scherrer equation: $D_{hkl} = K\lambda/(\beta \cos \theta)$ (*K*: Scherrer number (0.9), λ : incident ray wavelength (0.1542 nm), β : peak width at half height (rad), θ ; Bragg angle). ²⁹Si cross polarization magic angle spinning nuclear magnetic resonance $(^{29}Si$ CP-MAS NMR) measurements were obtained by an Avance III 500 (Bruker Analytik GmbH) in a 4 mm $ZrO₂$ rotor. The spinning rate was 8 kHz. The 29 Si chemical shifts are referenced to hexamethylcyclotrisiloxane (taken to be at $\delta = -9.6875$ ppm). Transmission electron microscope – energy dispersive X-ray spectroscopy (TEM-EDS) elemental mapping analytical techniques were done with a JEM-ARM200F (JEOL) at 200 kV. Inductively coupled plasma – atomic emission spectrometry (ICP-AES) was operated by an iCAP 6300 Duo (Thermo Fisher Scientific Inc.) to estimate the actual amount of precipitated $M(OH)_x$ and SiO_2 in asprepared $SiO_2@Mg-Al$ LDHs with various $Si/(Mg + Al)$ atomic ratio.

Results and discussion

Optimization of $Si/(Mg + Al)$ ratio in $SiO_2@LDH$

We prepared $SiO₂(QLDHs with various loading amounts of $SiO₂$$ to ascertain the optimized $SiO_2@LDH$ structure for high catalytic reactivity. The sphere morphology and the diameter of $SiO₂$ were confirmed from SEM and TEM observations, as presented in an earlier report.⁴⁷ The correlation between $Si/(Mg + Al)$ atomic ratio, base catalysis, and structural properties of $SiO₂(QLD)$ were investigated in the range of 0–0.50 on Si/(Mg +

Fig. 2 XRD patterns of as-prepared $SiO₂GLDHs$ with various $Si/(Mq +$ Al) atomic ratio.

Table 2 Crystal properties of as-prepared $SiO_2@LDHs$ with various $Si/(Mg + Al)$ atomic ratio

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	Table 2 Crystal properties of as-prepared $SiO2@LDHs$ with various $Si/(Mg + Al)$ atomic ratio			
$Si/(Mg + Al)$	Lattice	Crystallite size	Lattice	Crystallite
atomic ratio	parameter c/nm	$(003)^{a}/nm$	parameter <i>a</i> /nm	size $(110)^{a}/nm$
$\bf{0}$	2.33	7.6	0.31	13.4
0.13	2.32	5.2	0.31	13.7
0.17	2.36	4.5	0.31	10.3
0.25	2.38	3.7	0.31	8.0
0.50	2.43	2.8	0.31	8.3
	Al) atomic ratio. The prepared catalysts are designated as $SiO_2@(Z)$ LDH, where Z is a $Si/(Mg + Al)$ atomic ratio. The actual		$SiO2$ surface through the Si-O-Al and Si-O-Mg bonds to inhibit ab-face stacking without reducing the plane crystallite size. This	
are presented in Table 1.	ratios of Si/ $(M^{2+} + M^{3+})$ and M^{2+}/M^{3+} in the obtained materials Fig. 1 presents catalytic activity for the Knoevenagel condensation over $SiO_2@(Z)$ LDHs as (A) time-based reaction progression on benzaldehyde conversion and (B) reaction rate. The detailed results are listed in Table S1 (see ESI†). Among various $\frac{Si}{Mg} + Al$ atomic ratio from 0 to 0.50, actually, the 0.17 was found to be the best catalyst with a reaction rate of 171.1 mmol $g(cat)^{-1} h^{-1}$. This reaction rate is 2.2 times higher than conventional LDH prepared with the same co-precipitation method without SiO_2 seeds $(Si/(Mg + Al) = 0)$. The XRD patterns and crystal properties of $SiO_2@(Z)$ LDHs		result seems to be attributable to the lower number of starting points of LDH crystal growth: the amount of metal constituting one crystal did not change compared with conventional LDH prepared without SiO_2 seeds $(Si/(Mg + Al) = 0)$. Actually, the proportion for the ²⁹ Si CP-MAS NMR peaks attributed to Si-O- Mg and Si-O-Al bonds on the $SiO_2@(0.13)$ LDH is only $\leq 40\%$, whereas that of $SiO_2(\mathbb{Q}(0.17)$ LDH is $\leq 61\%$, as shown in Fig. 3 and Table 3. Therefore, we infer that the number of Si-O-Mg and Si-O-Al bonds on the SiO ₂ surface deeply affected	

The XRD patterns and crystal properties of $SiO₂(Q/Z)$ LDHs are portrayed respectively in Fig. 2 and Table 2. All prepared catalysts showed an LDH-originated diffraction pattern. The intensity of LDH originated peaks decreased in accordance with Si loading amount, whereas that of amorphous $SiO₂$ increased slightly. Lattice parameters a and c , respectively calculated from LDH (003) and (110) diffraction peaks, are almost identical among $\frac{Si}{Mg + Al}$ atomic ratios of 0-0.50. This result indicates clearly that these $SiO_2(\mathfrak{D}(Z)$ LDHs have the same LDH crystal unit. However, the crystallite size of LDH is unquestionably affected by $Si/(Mg + Al)$ atomic ratio. The crystallite size of $D(003)$ is reduced in accordance with the Si loading amount. The crystallite size of $D(110)$ is almost identical in the region among $Si/(Mg + Al)$ ratio of 0–0.13, but it is reduced from *ca*. 13 nm to 8 nm when a $\frac{Si/(Mg + Al)}{atio}$ increased. In our earlier research, it was revealed that the co-precipitation method with co-existence of spherical $SiO₂$ caused dispersion of starting points of LDH crystal growth on the $SiO₂$ surface through the Si-O-Al and Si-O-Mg covalent bonds to lead generation of finecrystallized LDH nanocrystal. 47 Fig. 3(A) shows that the spherical SiO_2 (40 nm) showed three peaks at -91 , -100 and -109 ppm, which respectively correspond to Q^2 , Q^3 , and Q^4 species⁴⁸⁻⁵⁰ where $Qⁿ$ designated the Si-centered tetrahedral structural species; Q refers to silicon atom and n denotes the number of bridging oxygens. Furthermore, $SiO_2@(Z)$ LDHs showed broad resonance between -70 to -115 ppm, which include some peaks attributed to Q^0 and/or Q^1 (–60 to –83 ppm)⁴⁹ and a Si-centered tetrahedral structure that possesses Si–O–Al and Si–O–Mg bonds $(-73 \text{ to } -105 \text{ ppm}).^{42}$

These results suggest that, in the case of lower $Si/(Mg + Al)$ atomic ratio (<0.13), the LDH crystal is immobilized onto the

Fig. 3 ²⁹Si CP-MAS NMR spectra of (A) spherical SiO₂ (40 nm), (B) $SiO_2@(0.13)$ LDH, (C) $SiO_2@(0.17)$ LDH and (D) $SiO_2@(0.50)$ LHD.

Table 3 Surface area and silicon environments in the $SiO₂$ and $SiO₂@$ (Z)LDHs as determined by ²⁹Si CP-MAS NMR

Sample	Assignment	δ /ppm	Percentage/%	per obtained LDH phase and TOFbase were also maximized at Si, $(Mg + Al)$ ratio of 0.17 with the reaction rate of 193.6 mmo.
$SiO2$ (40 nm)	Q^4	-109	11	$g(LDH)^{-1}$ h ⁻¹ and TOF _{base} of 450 h ⁻¹ . Above the Si/(Mg + Al
sphere	Q^3	-100	70	ratio of 0.17, both the reaction rate per LDH phase and TOFbase
		-91	19	were decreased respectively to 158.4 mmol $g(LDH)^{-1}$ h ⁻¹ and
SiO ₂ (0.13)LDH	\tilde{Q}^3	-99	17	238 h ⁻¹ at Si/(Mg + Al) ratio of 0.50. These results strongly
	Q^2	-92	10	
	$Q^4(3Al)$, $Q^3(1Mg)$ $Q^4(4Al)$, $Q^3(1Al)$, $Q^2(1Mg)$	-85	30	suggest that the $Si/(Mg + Al)$ atomic ratio affects not only the LDH crystallite size and base amount but also the type of base
	\mathbf{Q}^1 and/or \mathbf{Q}^0	-79	43	sites and these fractions. ⁵¹
SiO ₂ (0.17)LDH	Q^3	-100	18	The LDH crystallite size of $SiO_2@(0.50)$ LDH is at least
	Q^2	-92	17	smaller than that of $SiO_2@(0.17)$ LDH. Therefore, the base
	$Q^4(3Al)$, $Q^3(1Mg)$			catalysis of $SiO_2@(0.50)$ LDH is expected to be better than that of
	$Q^4(4Al)$, $Q^3(1Al)$,	-85	44	$SiO2(0.17)$ LDH if the base catalysis is only influenced by the
	$Q^2(1Mg)$			crystallite size. ²⁹ Si CP-MAS NMR spectra showed that the
	Q^1 and/or Q^0	-78	21	proportion of terminal Si-OH species assigned as Q^0 and/or Q^3
SiO ₂ (0.50)LDH	Q^3	-101	14	
	O^2	-92	31	decreased in accordance with $Si/(Mg + Al)$ atomic ratio, as
	$Q^4(3Al)$, $Q^3(1Mg)$			shown in Table 3, indicating first that a surface Si-O-Si bond is
	$Q^4(4Al)$, $Q^3(1Al)$,	-85	37	cleaved to generate terminal Si-OH species and then that these
	$Q^2(1Mg)$			act as cross-link point with Mg and Al ions. Consequently, wher
	\mathbf{Q}^1 and/or \mathbf{Q}^0	-78	17	there are the excess free terminal Si-OH species in the solution
				after the generation of SiO ₂ @LDH, these excess Si-OH species
				cover the LDH crystal to produce Si-O-Mg and Si-O-Al covalent
	reduction of the crystallite size not only in the stacking direc-			bonds. Although the base amount is increased even the regior
	tion but also in the plane direction when below $Si/(Mg + Al)$ <			from $SiO_2@(0.13)$ LDH to $SiO_2@(0.50)$ LDH, this phenomenor
$0.17 - 0.25$.				might take place only with difficulty on the inferior base sites
	Correlation between the base amount and catalytic activity			located at a flat plane of LDH. However, the decrease of TOFbase
	for Knoevenagel condensation over $SiO_2@(Z)$ LDHs is presented			strongly suggests that the high active base sites are poisoned by
	in Table 4. Although the base amount of $SiO_2(\mathfrak{D}(0.13)$ LDH was			Si species in the case of a higher $Si/(Mg + Al)$ atomic ratio.
	lower than that of conventional LDH, the reaction rate and the			Dark-field TEM images and results of EDS elemental
	apparent TOF per base site (TOF _{base}) for $SiO_2@(0.13)$ LDH are			mapping of $SiO2(Q/Z)$ LDHs are presented in Fig. 4. In the case of
	higher than those of LDH. The $D(003)$ of $SiO2(0.13)$ LDH was			lower $Si/(Mg + Al)$ atomic ratio such as 0.13 and 0.17, the LDH
smaller than LDH, whereas $D(110)$ of $SiO2(0.13)$ LDH and LDH				
				crystal is generated with covering the $SiO2$ phase to form a $SiO2$

Correlation between the base amount and catalytic activity for Knoevenagel condensation over $SiO_2@(Z)$ LDHs is presented in Table 4. Although the base amount of $SiO₂(0.13)$ LDH was lower than that of conventional LDH, the reaction rate and the apparent TOF per base site (TOF_{base}) for $SiO_2(\mathfrak{D}(0.13)$ LDH are higher than those of LDH. The $D(003)$ of $SiO₂(0.13)$ LDH was smaller than LDH, whereas $D(110)$ of SiO₂@(0.13)LDH and LDH are almost identical, as shown in Table 2. Therefore, these indicated that the immobilization of LDH crystal onto $SiO₂$ with inhibition of the *ab*-face stacking led to increase in the number of highly active base sites located on the surface LDH layer. Above the $Si/(Mg + Al)$ atomic ratio of 0.13, a base amount increased in accordance with $Si/(Mg + Al)$ ratio from 0.32 to 0.49 mmol $\rm g (cat)^{-1}.$ Furthermore, the activity was maximized at

Dark-field TEM images and results of EDS elemental mapping of $SiO_2@(Z)$ LDHs are presented in Fig. 4. In the case of lower $Si/(Mg + Al)$ atomic ratio such as 0.13 and 0.17, the LDH crystal is generated with covering the $SiO₂$ phase to form a $SiO₂$ core – LDH shell-like structure, as shown in Fig. $4(A)$ – (I) . Furthermore, results show that the boundary between $SiO₂$ phase and LDH phase becomes ambiguous in accordance with the increase of $Si/(Mg + Al)$ atomic ratio (Fig. 4(K)–(T)). These indicate that first the LDH crystal grows up from the $SiO₂$ surface to generate the immobilized $SiO₂$ core - LDH shell structure. If there are excess dissolved $SiO₂$ species possessing

^a The amount of LDH in the SiO₂@(Z)LDHs was calculated by ICP-AES with an assumption: all SiO₂@(Z)LDHs are composed of mixture of LDH and SiO₂. ^b Reaction rate for the Knoevenagel condensation of benzaldehyde with ethyl cyanoacetate. \overline{c} Base amount calculated from poisoning test by benzoic acid titration.

Fig. 4 (A, F, K and P) Dark-field TEM images of as-prepared SiO2@LDHs with various Si/(Mg + Al) atomic ratio: (A) 0.13, (F) 0.17, (K) 0.25 and (P) 0.50. Also shown are (B–E), (G–J), (L–O), (Q–T) EDS mapping results of as-prepared SiO₂@LDH with various Si/(Mg + Al) atomic ratio.

a free terminal Si–OH group, then these produced Si–O–Mg and Si–O–Al covalent bonds with the LDH crystal to cover the LDH shell.

Accordingly, we infer the correlation between $Si/(Mg + Al)$ atomic ratio and base catalysis of prepared $SiO₂(\mathfrak{D}/\mathfrak{D})$ as follows: (i) below $Si/(Mg + Al)$ atomic ratio of 0.13, the LDH crystal is just immobilized onto the $SiO₂$ surface with inhibition of the ab-face stacking. The exposed corner and edge located at the surface layer act as highly active base sites. (ii) Fine crystallization occurs not only in the stacking direction, but also in the plane direction to increase the amount of base sites, especially base sites with high activity up to $Si/(Mg + Al)$ atomic ratio of 0.17. (iii) Above a $\mathrm{Si}/(\mathrm{Mg} + \mathrm{Al})$ atomic ratio of 0.17, excess terminal Si–OH species covered the highly active base sites to produce Si–O–Mg and Si–O–Al covalent bonds, thereby lowering the activity.

Conclusions

The effects of $SiO₂$ loading amounts on the crystallite sizes, basicity, and catalytic activity of $SiO₂(QLDH)$ catalysts were investigated. $SiO₂(QLDHs)$ were prepared using co-precipitation with the coexistence of various amounts of spherical $SiO₂$ with particle sizes of ca. 40 nm. The XRD results suggest that the LDH crystallite size of $D(003)$ is simply reduced in accordance with the Si loading amount. Furthermore, the crystallite size of $D(110)$ is almost identical below Si/(Mg + Al) of 0.13, although it is reduced above 0.13. Base catalysis of $SiO₂(QLD)$ was evaluated using Knoevenagel condensation of benzaldehyde and ethyl cyanoacetate. Both the reaction rate and the apparent TOF per base site were increased in the region between $Si/(Mg + Al)$ atomic ratio of 0–0.17, whereas the base amount is increased linearly from the $\frac{Si}{Mg + Al}$ atomic ratio of 0.13 to 0.50 with reduction of the LDH crystallite size. The results of ²⁹Si CP-MAS NMR and STEM-EDS suggest that a surface Si–O–Si bond is cleaved, generating terminal Si–OH species that act as a crosslink point with Mg and Al ions to form an immobilized $SiO₂$ core – LDH shell structure. However, when the amount of Si becomes excessive with respect to Mg and Al ions, the excess Si– OH group forms Si–O–Mg and Si–O–Al covalent bonds with LDH crystal to cover the LDH shell. From these results, we inferred the effect of $SiO₂$ amount on heterogeneous base

catalysis of SiO_2 @Mg–Al LDH as follows: (i) below $Si/(Mg + Al)$ atomic ratio of 0.13, the highly active base sites located at the corner and edge of surface layer are exposed by immobilization of LDH crystals onto the SiO_2 surface, (ii) up to $Si/(Mg + Al)$ atomic ratio of 0.17, the number of exposed highly active base sites is increased in accordance with reduction of LDH crystallite, and (iii) above $\frac{Si}{Mg} + Al$ atomic ratio of 0.17. Excess terminal Si–OH species covered the highly active base sites through the Si–O–Mg and Si–O–Al covalent bonds to decrease the activity. This study elucidated the correlation between $SiO₂$ amount, crystal properties, and the basicity of fine-crystallized $SiO₂@LDH$ catalysts to present a new technique to improve the base catalysis of the widely used LDH material. After optimization of the Si/(Mg + Al) atomic ratio in SiO₂@LDHs, the Si/ $(Mg + Al)$ atomic ratio of 0.17 was found to be the best catalyst, with the reaction rate of 171.1 mmol $\rm g (cat)^{-1} \, h^{-1},$ which is a 2.2 times higher value than that of the conventional LDH prepared with same protocol in absence of $SiO₂$ seed agents. **BSC** Advances **Comparisting** Comparisting Comparisting Comparison Comparison Comparison Article is licensed under the second under the second under a state is licensed on 27. Downloaded on 27. Downloaded in the second un

Conflicts of interest

There are no conflicts to declare.

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