



Title	Elucidation of the local structure of active titanium(IV) sites on silica-based phase-boundary catalysts for alkene epoxidation with aqueous hydrogen peroxide
Author(s)	Ikeue, Keita; Ikeda, Shigeru; Watanabe, Atsushi; Ohtani, Bunsho
Citation	Physical Chemistry Chemical Physics, 6(9), 2523-2528 https://doi.org/10.1039/b316818b
Issue Date	2004
Doc URL	http://hdl.handle.net/2115/48673
Rights	Phys. Chem. Chem. Phys., 2004, 6, 2523-2528 - Reproduced by permission of the PCCP Owner Societies
Type	article (author version)
File Information	PCCP6_2523.pdf



[Instructions for use](#)

Bunsho Ohtani

Catalysis Research Center, Hokkaido University, Sapporo 001-0021, Japan

TEL: +81-11-706-9132 FAX: +81-706-9133

E-mail: ohtani@cat.hokudai.ac.jp

Elucidation of the local structure of active titanium(IV) sites on silica-based phase-boundary catalysts for alkene epoxidation with aqueous hydrogen peroxide

Keita Ikeue,[‡] Shigeru Ikeda,^{‡,†} Atsushi Watanabe,[§] Bunsho Ohtani^{*,§}

[‡] “Conversion and Control by Advanced Chemistry”

PRESTO, Japan Science and Technology Agency (JST), Japan

[†] Research Center for Solar Energy Chemistry, Osaka University, Toyonaka 560-8531, Japan

[§] Catalysis Research Center, Hokkaido University, Sapporo 001-0021, Japan

Structural and functional aspects of active titanium sites for phase boundary catalytic (PBC) epoxidation of 1-octene with hydrogen peroxide (H_2O_2) were investigated in detail using X-ray absorption fine structure (XAFS) analysis and diffuse reflectance (DR) spectroscopy. By analysis of the Ti K-edge X-ray absorption near edge fine structure (XANES) spectra of several titanium-loaded silica catalysts, the ratio of the amount of 4-coordinated titanium oxide (T_{tet}) to that of a 6-coordinated one (T_{oct}) was determined. Monotonic increments of catalytic activity for epoxide production and efficiency of H_2O_2 utilization with the ratio $\text{T}_{\text{tet}}/(\text{T}_{\text{tet}}+\text{T}_{\text{oct}})$ revealed that the highly active catalysts mainly include T_{tet} but not

T_{oct} . DR spectra of samples with external surfaces partially covered with alkylsilyl groups indicated that there are at least two kinds of T_{tet} sites with different ligands. One site anchors an acidic hydroxyl ($T_{\text{tet}}(\text{OH})$), giving absorption centered at the wavelength of ca. 230 nm and the other, exhibiting an absorption peak at the wavelength of ca. 210 nm, is directly attached to an alkylsilyl group ($T_{\text{tet}}(\text{OSiR})$) formed via reaction of an acidic hydroxyl of $T_{\text{tet}}(\text{OH})$ with an alkylsilane reagent. Since the catalysts have both an alkylsilyl-grafted hydrophobic surface and a hydroxyl-terminated hydrophilic surface, it is postulated that the former is located on both hydrophilic ($T_{\text{tet}}(\text{OH,w})$) and hydrophobic ($T_{\text{tet}}(\text{OH,o})$) surfaces and the latter, $T_{\text{tet}}(\text{OSiR})$, exists only on the hydrophobic surface. From analyses of catalytic activities of several catalysts with different distributions of these T_{tet} sites, both $T_{\text{tet}}(\text{OH,o})$ and $T_{\text{tet}}(\text{OSiR})$ sites on the hydrophobic surface were proved to be active, while $T_{\text{tet}}(\text{OH,w})$ on the hydrophilic surface did not work for the present PBC system. Among the sites on the hydrophobic surface, moreover, it was found that a $T_{\text{tet}}(\text{OSiR})$ site acted as a more effective site for selective epoxidation when compared with $T_{\text{tet}}(\text{OH,o})$, which induced ring-opening of epoxide to give a by-product, 1,2-octanediol.

Introduction

In the known chemical processes in catalytic oxidation, liquid-phase epoxidation of alkene is one of most important reactions for producing fine organic chemicals and their precursors.¹ From the viewpoint of the establishment of environmentally benign and ecologically more acceptable chemical processes, the use of aqueous hydrogen peroxide (H_2O_2) as an oxidizing agent is desirable because of its low toxicity and environmentally benign characteristics, i.e., only releasing water as a by-product. However, the use of H_2O_2 in liquid-phase epoxidation reactions sometimes requires the addition of a co-solvent and/or agitation of the mixture to make an emulsion for achieving effective interaction of H_2O_2 with catalysts and substrates because H_2O_2 is supplied as an aqueous solution and reaction substrates are generally immiscible with water.²⁻⁴

Recently, we have proposed a novel catalytic system, named phase-boundary catalysis (PBC), for driving the liquid-phase oxidation of water-immiscible organic compounds using aqueous H_2O_2 and a solid catalyst without the requirement of a co-solvent or agitation.^{5,6} We have reported that titanium oxide-loaded NaY zeolite particles with external surfaces partially covered with alkylsilyl groups were spontaneously assembled at the liquid-liquid phase boundary and efficiently catalyzed epoxidation of straight-chain alkenes at room temperature even without agitation. It has been also revealed through fluorescence microscopic observation of the NaY particles selectively modified with fluorescence dye that the attached alkylsilyl groups were distributed only on one side of the external surface of the NaY particle, resulting in a bimodal surface structure, i.e., one side of the external surface is hydrophilic and the other is hydrophobic.⁷ To the best of our knowledge, there have been no reported examples of a solid catalyst- H_2O_2 system similar to this PBC system using bimodal catalyst particles.

The phenomenological aspects of PBC have been reported in detail in our previous papers⁶ and the bimodal characteristics have also been discussed in detail,⁷ but the structure of active sites has remained unclear. It is necessary to clarify and control the local structure of active species in catalysts for further applications of PBC to various substrates. In the present study, therefore, we focused our attention on the local structure of titanium oxide species on support materials to obtain information that will be useful for the design and preparation of suitable bimodal particles for PBC.

Experimental

Silica (Degussa AEROSIL 200) loaded with various amounts of titanium(IV) oxide species ($C_{Ti} / \mu\text{mol g}^{-1}$) was prepared by the following procedure. Three grams of silica was added to 100 cm³ of 2-propanol solution of titanium(IV) tetraethoxide ($\text{Ti}(\text{OEt})_4$) under a dry argon atmosphere. The amount of $\text{Ti}(\text{OEt})_4$ was adjusted in order to synthesize samples with different C_{Ti} . After stirring the suspension at room temperature for 3 h, a solid part was collected by centrifugation, dried at 383 K for 2 h under vacuum, and calcined at 773 K for 6 h. The as-prepared samples are called w-Ti(C_{Ti})/SiO₂, e.g., w-Ti(44)/SiO₂ represents a sample containing 44 $\mu\text{mol g}^{-1}$ of titanium atom. The titanium content (C_{Ti}) was determined by colorimetric analysis of the samples using 1,2-dihydroxy-3,5-benzenedisulfonic acid, disodium salt (Tiron) as a colorimetric agent. Bimodal particles whose external surfaces were partially covered with alkylsilyl groups were prepared as follows. To 10 cm³ of toluene containing 1 mmol of n-octadecyltrimethoxysilane (ODMS), 0.5 g of w-Ti(C_{Ti})/SiO₂ thoroughly mixed with 16.7 wt% of water was added, and the suspension was shaken for 2 min at room temperature. After the addition of triethylamine (1 mmol), the suspension was further shaken for 10 min at room temperature. Then the solid was collected by centrifugation and dried at 383 K for

2 h under vacuum. The thus-prepared samples were labeled w/o-Ti(C_{Ti})/SiO₂. A sample whose external surfaces were covered with a silica shell was also prepared by the reaction of dry w-Ti(158)/SiO₂ and ODMS followed by calcination at 773 K for 6 h to remove the organic moieties.

In a typical experiment for epoxidation of 1-octene with aqueous hydrogen peroxide, 1-octene (4 cm³), 30% aqueous hydrogen peroxide (1 cm³) and catalyst particles (50 mg) were placed in a glass tube (16 mm in diameter) and kept at 298 K for 20 h. In order to remove appreciable impurities, 1-octene (Kanto Chemical) was treated with basic alumina (Merck) before use. Reaction products were analyzed by a Shimadzu GC-14B gas chromatograph equipped with a DB-1 column (J&W) and an FID detector.

Diffuse reflection (DR) spectra were recorded on a Hamamatsu Photonics C7473-6 Photonic Multi-Channel Analyzer, and the reflection data were converted to Kubelka-Munk (KM) function. Before the DR measurements, all samples were dried at 573 K or 423 K for 3 h under vacuum and the data were collected at room temperature under vacuum. XAFS measurements were carried out at the BL-9A facility⁸ of the Photon Factory at High Energy Acceleration for Research Organization, Tsukuba, Japan (Proposal #2002G272). The X-ray source from a 2.5 GeV electron-storage ring was monochromatized using an Si(111) double-crystal monochromator. The Ti K-edge absorption spectra were recorded in the fluorescence mode at 298 K after pretreatment of the samples at 573 K for 3 h under vacuum. The XANES spectra were analyzed using a Rigaku REX 2000 program.

Results and Discussion

Catalytic activity for epoxidation of 1-octene.

In the case of NaY zeolite, which was used as a support in our previous studies,⁵⁻⁷ several factors such as pore structure and acid property might influence both structural characteristics of active sites and catalytic properties. Therefore, in order to determine the pure effects of local structure of active sites on reactivity and selectivity, nonporous silica was selected as a model support material. Table 1 summarizes the yields of products from the mixture of 1-octene and aqueous H₂O₂ in the presence of titanium-loaded silica samples. Upon dispersion in the two-phase mixture, w/o-Ti(C_{Ti})/SiO₂ was spontaneously assembled at the liquid-liquid phase boundary (Fig. 1a) and showed activity for epoxidation to give 1,2-epoxyoctane as a main or a sole product even without agitation (entries 1 and 2). Note that, probably due to the successive ring-opening reaction (see below) with stirring, w/o-Ti(333)/SiO₂ with a relatively high titanium content gave less than half yield of epoxide with stirring (entry 2).⁶ On the other hand, the w-Ti(158)/SiO₂ sample, which preferentially disperses in the aqueous phase as shown in Fig. 1b, showed an appreciable, but quite low, level of activity even with stirring (entry 3). Since these trends are similar to the previous results obtained by using titanium(IV) oxide-loaded NaY zeolite catalysts, the present w/o-Ti(C_{Ti})/SiO₂-based system is also an example of phase-boundary catalysis, though details of the distribution of surface-covering alkylsilyl groups and the bimodal unsymmetrical surface characteristics of silica particles were not discussed here.⁹ Since it seems that w/o-Ti(C_{Ti})/SiO₂ does not require mechanical agitation, the catalytic reactions reported in this paper were carried out without agitation unless otherwise stated.

Table 2 summarizes dependence of the catalytic activity of w/o-Ti(C_{Ti})/SiO₂ for 1-octene epoxidation on C_{Ti}. For samples with relatively low C_{Ti} (entries 1-3), 1,2-epoxyoctane was produced as a sole product and its yield increased with increase in C_{Ti}, while by-products, e.g., 1,2-octanediol, appeared in reaction with catalysts containing a relatively large amount of titanium (entries 4 and 5) and the

epoxide yield was reduced along with C_{Ti} . Moreover, turnover number (TON) based on molar amount of titanium atom and efficiency of H_2O_2 utilization ($E_{\text{TH}_2\text{O}_2}$), defined as percent proportion of H_2O_2 used for epoxide production among the total consumption of H_2O_2 , decreased gradually with increase in C_{Ti} . Since structure and dispersibility of Ti on silica surfaces may depend on its content, the observed C_{Ti} dependence might be induced by such structural alteration of titanium sites. Thus, the difference in local structure of active sites between these catalysts will be discussed in the following sections.

Correlation between coordination number of Ti sites and catalytic activity.

Figure 2 shows typical DR spectra of w-Ti(C_{Ti})/SiO₂ with different values of C_{Ti} . All of the samples gave absorption in the ultraviolet region below ca. 300 nm with a peak at ca. 230-240 nm. Since the original silica has no photoabsorption in this wavelength range and, qualitatively, the intensity of absorption seems to increase with increase in Ti content (Fig. 2a), the absorption in all samples should be assigned to loaded titanium species. By a comparison of shapes of the absorption spectra normalized by KM function at 230 nm (Fig. 2b), it was revealed that these spectra tended to expand to a longer wavelength range with increase in C_{Ti} . According to the literature,¹⁰⁻¹³ absorption at a wavelength around 200-250 nm is assigned to ligand-to-metal charge transfer (LMCT) from oxygen atoms to Ti^{4+} of 4-coordinated titanium oxide species (T_{tet}), and absorption in a longer wavelength region (ca. 250-290 nm) is attributed to aggregated 6-coordinated titanium oxide species (T_{oct}). Therefore, it is presumed that the titanium sites on the present samples were a mixture of T_{tet} and T_{oct} and that their ratio depended strongly on the Ti content (see below).

Figure 3 shows the Ti K-edge XANES spectra of selected w-Ti(C_{Ti})/SiO₂ samples. Spectra of crystalline titanium(IV) oxide (TiO_2 , Degussa P25) and TS-1

zeolite are also shown (Figs. 3a and 3b) as typical examples of T_{tet} and T_{oct} , respectively. The XANES spectra have several well-defined peaks; TS-1, which includes mainly T_{tet} , exhibited an intense single preedge peak at 4969.5 eV, while three characteristic weak preedge peaks appeared for bulk TiO_2 composed of T_{oct} . As is expected from above results of the above-described DR spectra, $w\text{-Ti}(C_{\text{Ti}})/\text{SiO}_2$ showed spectral characteristics of both T_{tet} and T_{oct} . In order to quantify T_{tet} and T_{oct} in the samples, the ratio of T_{tet} to T_{oct} are determined by curve fitting of the XANES spectra of the samples to simulated spectra obtained by the linear combination of spectra of TS-1 (T_{tet}) and TiO_2 (T_{oct}). Their best fitted curves are shown in Fig. 3 (dotted lines), and thus-obtained ratios of T_{tet} to T_{tet} and T_{oct} ($T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$) are summarized in Table 2. It is reasonable that $T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$ is almost reciprocal to C_{Ti} ; $w\text{-Ti}(44)/\text{SiO}_2$, having the lowest C_{Ti} , included mainly T_{tet} , but higher loading of Ti induced aggregation between Ti atoms and thereby the $T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$ halved in samples with the highest Ti content ($w\text{-Ti}(528)/\text{SiO}_2$).

Figure 4 shows plots of TON and $E_{\text{H}_2\text{O}_2}$ as a function of $T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$. Both TON and $E_{\text{H}_2\text{O}_2}$ monotonically increased with increase in $T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$. Except for side reactions, e.g., ring-opening of epoxide (see below), the main chemical events in the present system are assumed to be (1) formation of epoxide from 1-octene with H_2O_2 and (2) decomposition of H_2O_2 presumably into molecular oxygen (O_2). Although it is practically impossible to analyze such events on titanium sites, T_{tet} and T_{oct} , independently, the present results suggest that T_{tet} mainly works as active sites for epoxide formation and that T_{oct} is relatively more active for the decomposition of H_2O_2 .

It should be noted that there was no appreciable activity on the hydrophilic $w\text{-Ti}(158)/\text{SiO}_2$ catalyst with low C_{Ti} that included T_{tet} as a main component of Ti species even though thorough mechanical agitation to drive mass transfer was provided (Table 1, entry 3). Therefore, for the use of phase-boundary catalytic

systems, additional structural features of active site(s), e.g., effective location of titanium sites on the surface of w/o-Ti(C_{Ti})/SiO₂, having both hydrophobic and hydrophilic faces, and local environment of T_{tet} after modification with alkylsilyl groups, should also be taken in consideration.

Role of Alkylsilyl groups in selective production of epoxide.

To obtain further information on the structure of active sites on the catalysts after modification with alkylsilyl groups (w/o-Ti(C_{Ti})/SiO₂), DR spectra were compared with the spectrum of w-Ti(C_{Ti})/SiO₂, i.e., the sample without modification with alkylsilyl groups. Figure 5 shows DR spectra of samples of C_{Ti} of 158 with and without modification. In order to avoid the collapse of surface alkylsilyl groups on w/o-Ti(158)/SiO₂, both spectra were obtained after pretreatment at relatively low temperature (423 K) in comparison with the spectra shown in Fig. 2, those of which were obtained after pretreatment at 573 K. However, based on the facts that there is no difference in the spectra of w-Ti(158)/SiO₂ between these pretreatment conditions, differences of water adsorption and/or hydration level, if present, should not influence the DR spectra. It is clear from this figure that modification results in no appreciable broadening and/or shift of the spectrum to a longer wavelength range, indicating that the coordination number of titanium species was not affected by the modification, i.e., aggregation between isolated titanium species (T_{tet}) was not induced, and T_{tet}/(T_{tet}+T_{oct}) might therefore not be changed by the modification.¹⁴ One of the most notable features shown in Fig. 5 is that the w/o-Ti(158)/SiO₂ gave two absorption peaks centered at the wavelengths of ca. 210 nm (A₂₁₀) and ca. 230 nm (A₂₃₀). Such absorption peaks have also been observed in dehydrated TS-1 and attributed to two different framework sites as [Ti(OH)(OSi)₃] and [Ti(OSi)₄],¹⁵ though their assignment to two peaks has not been fixed yet. Moreover, as has been reported by Machese et al., there were two kinds of

tetrahedral titanium sites in Ti-MCM-41 that were detected by photoluminescence measurements.¹⁶ Therefore, the two peaks in Fig. 5 are attributable to titanium sites in different tetrahedral environments. Based on the fact that A_{230} is similar to the predominant peak of the w-Ti(158)/SiO₂ sample with no alkylsilyl groups (Fig. 5a), the peak may correspond to the tetrahedral titanium site having a hydroxyl group (T_{tet}(OH)) (Fig. 6a, left (i)). On the other hand, since A_{210} appeared also in the spectrum of the silica-coated w-Ti(158)/SiO₂ sample, i.e., this peak seemed to appear only when a Ti unit is covered by -O-Si-O- shells, A_{210} is attributable to the species formed via reaction of a hydroxyl group on the tetrahedral titanium site (Ti-OH) with ODMS (T_{tet}(OSiR), see Fig. 6b, right (ii)).

In order to represent the distribution of these T_{tet} sites shown in Fig. 6a semi-quantitatively, the ratio of KM function at 210 nm to that at 230 nm (KM_{210}/KM_{230}) was introduced for convenience. Table 3 summarizes the calculated KM_{210}/KM_{230} values of the w/o-Ti(C_{Ti})/SiO₂ and w-Ti(158)/SiO₂ samples together with the selectivity of 1-octene epoxidation, defined as the percentage of epoxide yield in total amount of products. In order to exclude the influence of aggregated titanium species (T_{oct}) on the activity (Table 2), samples with ca. 120-160 $\mu\text{mol g}^{-1}$ of C_{Ti}, giving similar $T_{\text{tet}}/(T_{\text{tet}}+T_{\text{oct}})$ in this table were selected. Compared to w-Ti(158)/SiO₂, i.e., without ODMS modification, modified samples showed increment in KM_{210}/KM_{230} up to almost unity, indicating the formation of T_{tet}(OSiR) sites due to the modification. It should be noted, however, that a few samples showed relatively small or almost no increment in KM_{210}/KM_{230} due to modification with alkylsilyl groups even though the same ODMS treatment was conducted (entries 3 and 4). Such differences in the KM_{210}/KM_{230} values could not be elucidated and controlled at present, probably because these differences were caused by subtle unexpected differences in experimental conditions, e.g., humidity and contaminants in solvents. Anyway, among the w/o-Ti(C_{Ti})/SiO₂ samples with

different values of $K_{M_{210}}/K_{M_{230}}$, the higher $K_{M_{210}}/K_{M_{230}}$ was, the higher was the product selectivity. These results indicate that the $T_{\text{tet}}(\text{OSiR})$ site was an active site for selective epoxide formation.

Local structure of active sites for PBC.

A possible model for the active titanium sites was considered on the basis of the above discussion, and the overall structures of them on the $w/o\text{-Ti}(\text{C}_{\text{Ti}})/\text{SiO}_2$ are illustrated in Fig. 6. Similar to previously reported NaY zeolite-based samples, $w/o\text{-Ti}(\text{C}_{\text{Ti}})/\text{SiO}_2$ might have both hydrophobic and hydrophilic faces. Thus, the loaded T_{tet} was categorized into two groups: one on the hydrophilic surface ($T_{\text{tet}}(\text{OH},w)$) and the other on the hydrophobic surface of silica. Moreover, the above-described results suggest that the T_{tet} on the hydrophobic surface could be divided into two groups of sites: sites with no binding with alkylsilyl groups ($T_{\text{tet}}(\text{OH},o)$) and sites directly bound to alkylsilyl groups ($T_{\text{tet}}(\text{OSiR})$). From the fact that there was no appreciable activity on $w\text{-Ti}(\text{C}_{\text{Ti}})/\text{SiO}_2$ containing only $T_{\text{tet}}(\text{OH},w)$ even with vigorous stirring (Table 1, entry 3), $T_{\text{tet}}(\text{OH},w)$ on $w/o\text{-Ti}(\text{C}_{\text{Ti}})/\text{SiO}_2$ samples is thought to be inactive, probably owing to their less accessibility and low adsorption affinity to the substrate (1-octene). On the other hand, $T_{\text{tet}}(\text{OH},o)$ on the hydrophobic surface of the present PBC ($w/o\text{-Ti}(\text{C}_{\text{Ti}})/\text{SiO}_2$) is thought to be active because appreciable activity was seen even in the absence of or with only a small amount of $T_{\text{tet}}(\text{OSiR})$ (Table 3, entry 4). Corma et al. reported that the presence of $\equiv\text{Ti-OH}$ groups in Ti-zeolite allows adsorption of the product epoxide due to its higher polarity than that of olefinic substrates, resulting in a ring opening of epoxide to diols.¹⁷ The $T_{\text{tet}}(\text{OH},o)$ site, therefore, could induce a ring-opening reaction to yield 1,2-octanediol. Due to the absence of such acidic hydroxyl in the neighborhood, $T_{\text{tet}}(\text{OSiR})$ on the hydrophobic surface is thought to be an active site for selective production of epoxides. It is also probable to induce such ring opening

reaction on the surface Si-OH. However, there was no production of the diol when the catalyst with high $T_{\text{tet}}(\text{OSiR})$ was used even though the presence of Si-OH. Moreover, the diol was not produced when the mixture of epoxides, hydrogen peroxide, and SiO_2 particles without loading of titanium was stirred at room temperature. Therefore, the diol liberation was only induced on Ti-OH in the present phase-boundary catalytic system.

Conclusions

The local structure of titanium oxide species on silica for phase-boundary catalytic epoxidation of 1-octene with aqueous hydrogen peroxide was investigated. It has been proved that 4-coordinated titanium oxide species directly attached to alkylsilane ($T_{\text{tet}}(\text{OSiR})$) on the hydrophobic face of the bimodal particle is an active site for selective production of 1,2-epoxyoctane but that two other kinds of titanium species are inactive or active but non-selective for epoxide formation, inducing ring opening to yield 1,2-octanediol. At present, some extent of titanium atom could be loaded on silica as the site of a promising structure, $T_{\text{tet}}(\text{OSiR})$, for selective production of epoxide. Therefore, further improvement for selective loading of the $T_{\text{tet}}(\text{OSiR})$ site is expected. Although, in the present study, we used 1-octene as a sole substrate and the observed activity was not so high, there should be several substrates that can be used for the present phase-boundary catalytic oxidation with H_2O_2 and, actually, we have already found some reactions could be catalyzed efficiently in this system. Thus, further investigations along these lines are under way.

Acknowledgements

The authors are grateful to Nippon Aerosil Co., Ltd. for supplying silica samples. Mr. Tetsuzo Habu and Mr. Kazuhiro Matsudaira (Technical Division of the

Catalysis Research Center, Hokkaido University) are acknowledged for their assistance in the construction of the XAFS cells and the reaction apparatus.

References

1. M. Dusi, T. Mallat and A. Baiker, *Catal. Rev. Sci. Eng.*, 2000, **42**, 213-278.
2. A. Bhaumik and R. Kumar, *J. Chem. Soc. Chem. Commun.*, 1995, 349-350.
3. T. Tatsumi, K. A. Koyano and N. Igarashi, *Chem. Commun.*, 1998, 325-326.
4. A. Bhaumik, P. Mukherjee and R. Kumar, *J. Catal.*, 1998, **178**, 101-107.
5. H. Nur, S. Ikeda and B. Ohtani, *Chem. Commun.*, 2000, 2235-2236.
6. H. Nur, S. Ikeda and B. Ohtani, *J. Catal.*, 2001, **204**, 402-408.
7. S. Ikeda, H. Nur, T. Sawadaishi, K. Ijro, M. Shimomura and B. Ohtani, *Langmuir*, 2001, **17**, 7976-7979.
8. M. Nomura and A. Koyama, *Nucl. Instrum. Meth. A*, 2001, **467-468**, 733-736.
9. S. Ikeda, K. Ikeue and B. Ohtani, to be published.
10. J. Jarupatrakom and T. D. Tilley, *J. Am. Chem. Soc.*, 2002, **124**, 8380-8388.
11. C. Lamberti, S. Bordiga, D. Arduino, A. Zecchina, F. Geobaldo, G. Spanò, F. Genoni, G. Petrini, A. Carati, F. Villain and G. Vlaic, *J. Phys. Chem. B*, 1998, **102**, 6382-6390.
12. M. Raimondi, E. Gianotti, L. Marchese, G. Martra, T. Maschmeyer, J. M. Seddon and S. Coluccia, *J. Phys. Chem. B*, 2000, **104**, 7102-7109.
13. K. Ikeue, H. Mukai, H. H. Yamashita, S. Inagaki and M. Anpo, *J. Synchrotron Rad.*, 2001, **8**, 640-642.
14. To exclude the influence of the coordination of water molecules onto Ttet, XAFS should be measured after pretreatment under vacuum at > 573 K. However, due to the presence of alkyl groups on w/o-Ti(C_{Ti})/SiO₂, such pretreatment at a high temperature might result in decomposition of them.

Therefore, the coordination number of titanium oxide species on w/o-Ti(C_{Ti})/SiO₂ could not be accurately determined by XAFS measurement.

15. L. Le-Noc, D. Trong-On, S. Solomykina, B. Echchahed, F. Beland, D. Cartier dit Moulin and L. Bonneviot, *Stud. Surf. Sci. Catal.*, 1996, **101**, 611-620.
16. L. Marchese, T. Maschmeyer, E. Gianotti, S. Coluccia and J. M. Thomas, *J. Phys. Chem. B*, 1997, **101**, 8836-8838.
17. A. Corma, M. Domine, J. A. Gaona, J. L. Jordá, M. T. Navarro, F. Rey, J. Pérez-Pariente, J. Tsuji, B. McCulloch and L. T. Nemeth, *Chem. Commun*, 1998, 2211-2212.

Figure captions

Fig. 1. Photographs of w/o-Ti(C_{Ti})/SiO₂ and w-Ti(C_{Ti})/SiO₂ samples in a 1-octene/aqueous H₂O₂ (30%) binary system.

Fig. 2. (a) Representative diffuse reflectance (DR) spectra of w-Ti(C_{Ti})/SiO₂ samples and (b) those normalized at 230 nm. Titanium contents (C_{Ti} / $\mu\text{mol g}^{-1}$) were (1) 75, (2) 158, and (3) 528. All spectra were measured after evacuation at 573 K for 3 h.

Fig. 3. Representative Ti K-edge XANES spectra of several titanium oxide-containing samples. (a) TiO₂ (Degussa P-25), (b) TS-1, (c) w-Ti(528)/SiO₂, (d) w-Ti(158)/SiO₂. Dotted lines represent simulated spectra obtained by linear combination of spectra (a) and (b). Before measurements, all samples were degassed by evacuation at 573 K for 3 h.

Fig. 4. Plots of TON and $E_{H_2O_2}$ as a function of $T_{tet}/(T_{tet}+T_{oct})$. Titanium contents (C_{Ti} / $\mu\text{mol g}^{-1}$) were (1) 44, (2) 75, (3) 158, (4) 333, and (5) 528.

Fig. 5. DR spectra of (a) w-Ti(158)/SiO₂ and (b) w/o-Ti(158)/SiO₂. All spectra were measured after evacuation at 423 K for 3 h.

Fig. 6. Schematic illustrations of (a) the local structure and (b) the distribution of tetrahedral titanium oxide species on w/o-Ti(C_{Ti})/SiO₂ samples. (i) $T_{tet}(\text{OSiR})$ sites located on the hydrophobic surface with an absorption peak at ca. 210 nm, (ii) $T_{tet}(\text{OH})$ sites located on both hydrophilic and hydrophobic surfaces ($T_{tet}(\text{OH,w})$ and $T_{tet}(\text{OH,o})$, respectively) giving absorption centered at ca. 230 nm.

Table 1. Effect of agitation on the yield of epoxide by titanium-loaded silica particles^a

entry	catalyst	yield / μmol					
		static ^b			stirring ^c		
		epoxide ^d	diol ^e	others ^f	epoxide ^d	diol ^e	others ^f
1	w/o-Ti(158)/SiO ₂	72.8	0 ^g	0 ^g	87.1	4.2	2.7
2	w/o-Ti(333)/SiO ₂	41.9	3.9	1.1	16.7	10.8	8.5
3	w-Ti(158)/SiO ₂	0.3	0 ^g	0.1	0.5	0 ^g	0.2

^aAll reactions were carried out at 298 K for 20 h with 1-octene (4 cm³), 30 % H₂O₂ (1 cm³) and catalyst (50 mg). ^bReaction was carried out under static condition. ^cReaction was carried out with vigorous mechanical agitation. ^d1,2-Epoxyoctane. ^e1,2-Octanediol. ^f1-Octanol and 2-octanol. ^gNot detected ($< 10^{-3}$ μmol).

Table 2. Dependence of activity for epoxidation of 1-octene with H₂O₂ on titanium contents in various w/o-Ti(C_{Ti})/SiO₂ samples^a

entry	catalyst	T _{tet} /(T _{tet} +T _{oct}) ^b	yield / μmol			TON	Ef _{H₂O₂} ^f / %
			epoxide ^c	diol ^d	others ^e		
1	w/o-Ti(44)/SiO ₂	0.95	27.5	0	0	12.6	53
2	w/o-Ti(75)/SiO ₂	0.89	36.6	0	0	9.7	46
3	w/o-Ti(158)/SiO ₂	0.81	72.8	0	0	9.2	33
4	w/o-Ti(333)/SiO ₂	0.57	41.9	3.9	1.1	2.5	14
5	w/o-Ti(528)/SiO ₂	0.55	39.7	6.5	1.6	1.5	5

^aReaction conditions are the same as those given for Table 1. ^bRatio of the content of 4-coordinated titanium oxide (T_{tet}) to the sum of T_{tet} and that of 6-coordinated one (T_{oct}). ^c1,2-Epoxyoctane. ^d1,2-Octanediol. ^e1-Octanol and 2-octanol. ^fEfficiency of H₂O₂ utilization defined as the percentage of the epoxide yield to amount of consumed and decomposed H₂O₂.

Table 3. Selectivity of epoxidation of 1-octene with H₂O₂ by w/o-Ti(C_{Ti})/SiO₂^a and ratio of Kubelka-Munk functions

entry	catalyst	yield / μmol			selectivity ^e / %	KM ₂₁₀ /KM ₂₃₀ ^f
		epoxide ^b	diol ^c	others ^d		
1	w/o-Ti(122)/SiO ₂	55.6	0	0	100	1.00
2	w/o-Ti(158)/SiO ₂	72.8	0	0	100	1.01
3	w/o-Ti(131)/SiO ₂	61.3	3.0	1.6	93	0.97
4	w/o-Ti(159)/SiO ₂	50.2	7.5	1.7	85	0.93
5	w-Ti(158)/SiO ₂	0.3	0	0.1	75	0.90

^aReaction conditions are the same as those given for Table 1. ^b1,2-Epoxyoctane. ^c1,2-Octanediol. ^d1-Octanol and 2-octanol ^eSelectivity defined as the percentage of the yield of epoxide to other by-products. ^fRatio of K-M function at 210 nm to that at 230 nm.

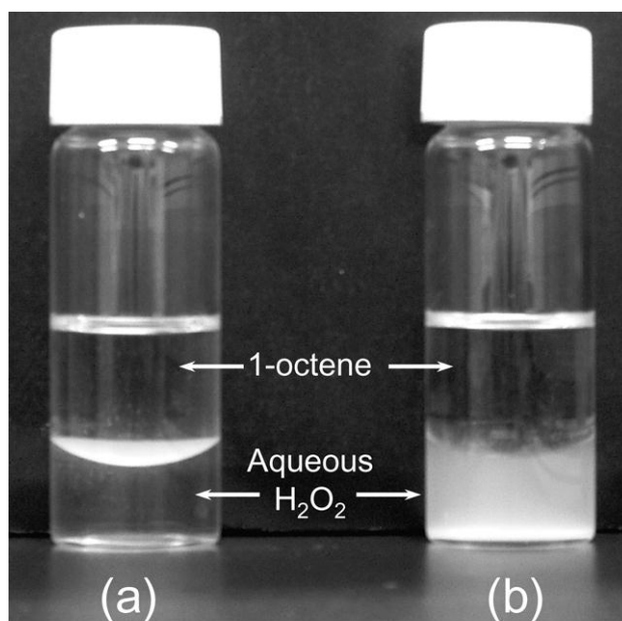


Figure 1. *K. Ikeue et al.*

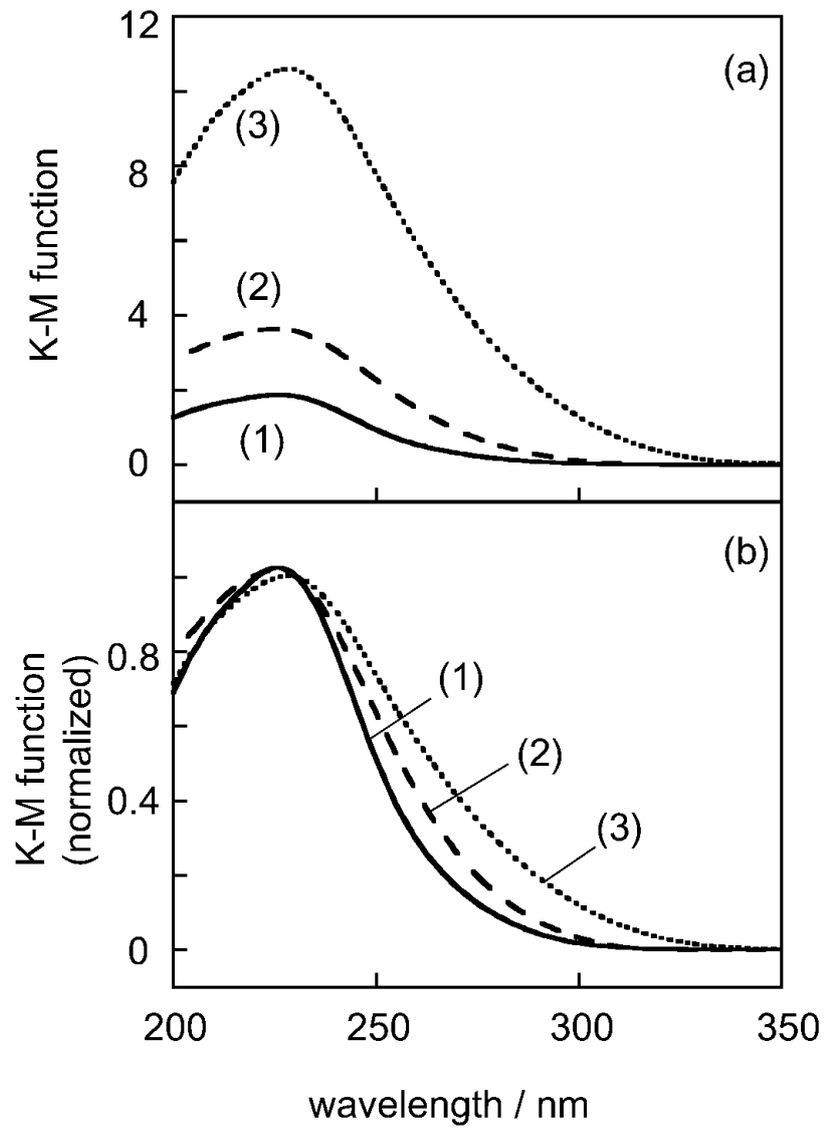


Figure 2. *K. Ikeue et al.*

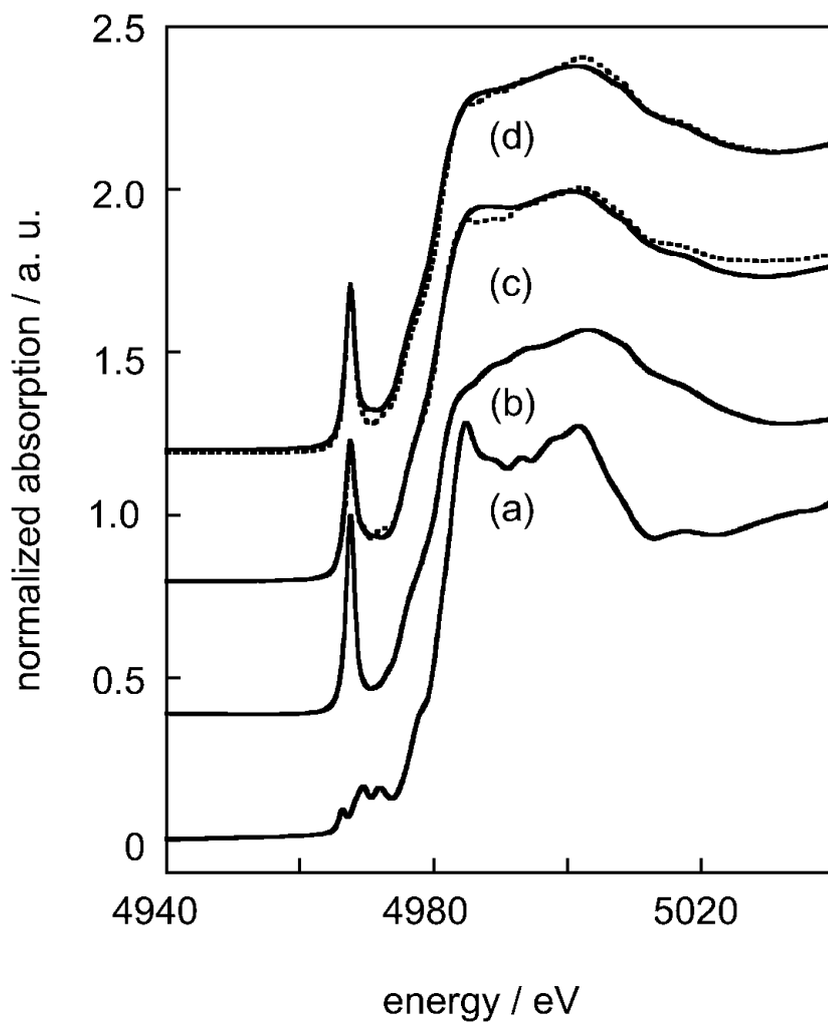


Figure 3. *K. Ikeue et al.*

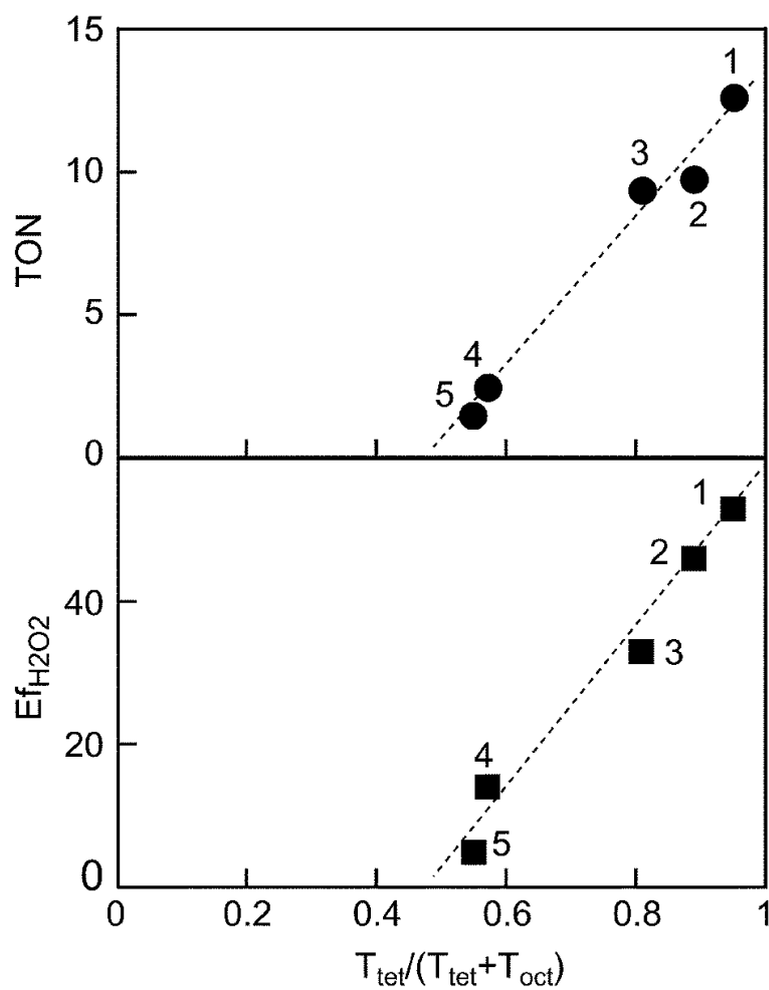


Figure 4. *K. Ikeue et al.*

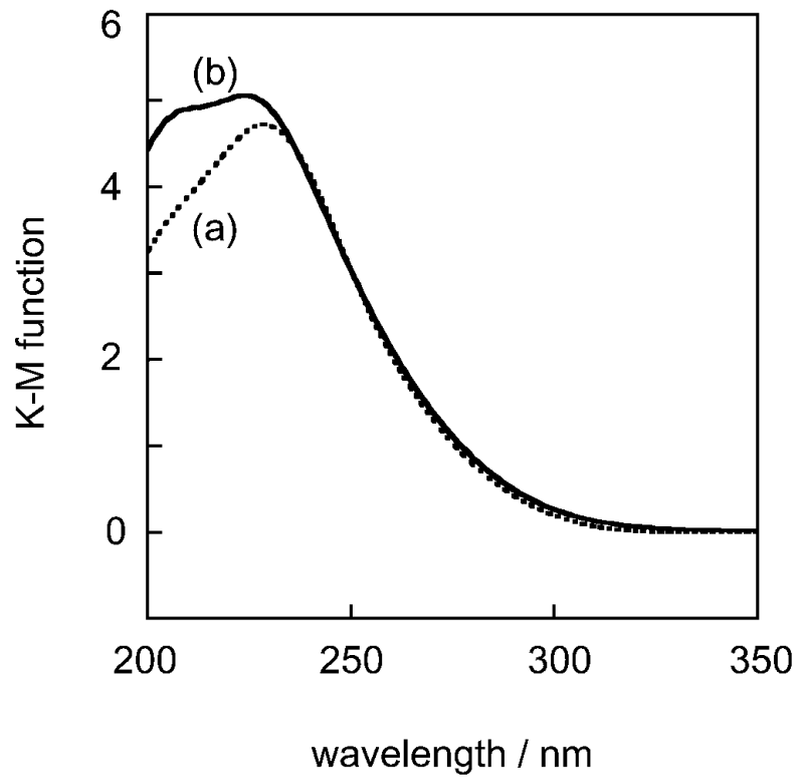


Figure 5. *K. Ikeue et al.*

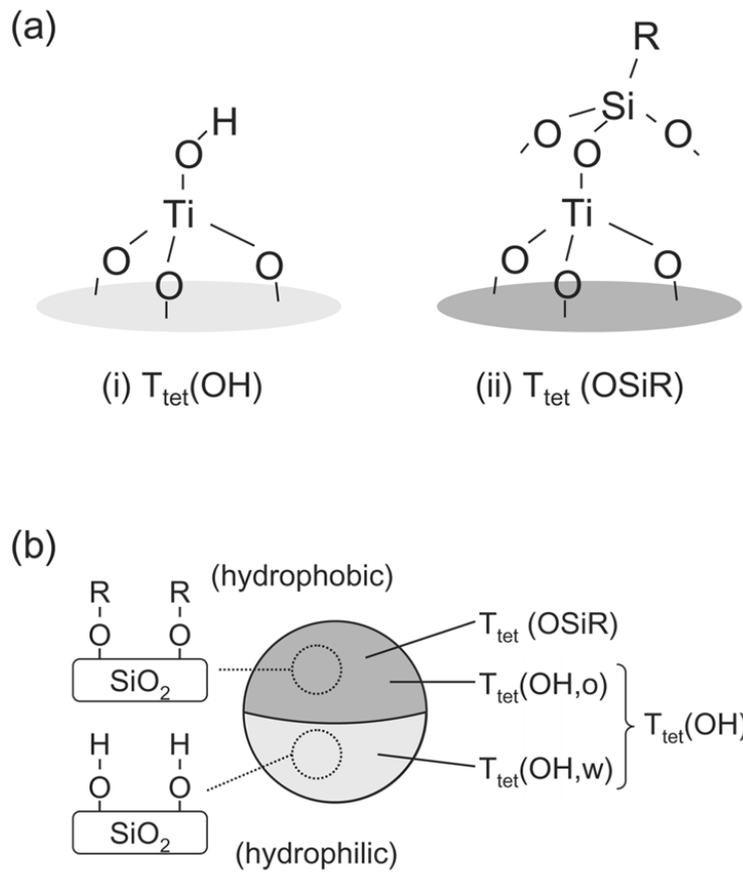


Figure 6. *K. Ikeue et al.*