Roles of interfaces in nanostructured composites: nanocatalysts, sponge crystals and thin films

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Interfaces are important in the functionality of materials with composite structures. Focusing mainly on our studies, this article provides an overview of the functionality that can be achieved by the careful design of interfaces in inorganic composites. Molecular selective photocatalysts have been realized by developing nanostructures, in which photocatalytically active metal oxide particles are surrounded by mesoporous silica. Such composites structure achieved combined function of molecular selective adsorption and photocatalytic decomposition. Solid acid catalysts that are active in water have been realized by exploiting the nanostructure of mesoporous silica. Catalytically active inorganic molecules such as heteropolyacids have been anchored on the walls within the pores of hydrophobic organic layers. The resulting nanocomposite exhibited high acid catalytic activity in water. Sponge crystals are a more recently discovered class of porous crystals. We discuss their structure, and propose revised classifications of microporous crystals, mesocrystals and sponge crystals. Thin films of metal nitrides such as Sr₂N and metal oxides have been prepared by molecular beam epitaxy and pulsed laser deposition. The effects of the interface between the film and substrate are discussed.

Key-words : Photocatalyst, Catalyst, Mesoporous silica, Sponge crystals, Nitride, Ferromagnetism

1. Introduction

Designing composite structures has for many years been an intensively studied field of materials science and chemistry. The nanoscale design of the structures of composite materials is a key strategy in various research fields. Composites contain two or more components or domains. As a consequence, they contain interfaces between these components.

A useful strategy for developing composites is to design interactions through these interfaces. To understand and design various functionality into composites, it is useful to consider what each side of an interface “exchanges”, “transfers”, or “communicates” through the interface. Components on either side of an interface in a composite can interact by:
- (i) transfer of molecules through the interface,
- (ii) transfer of electrons and/or holes through the interface,
- (iii) inducing electric carriers through a strong electric field at the interface,
- (iv) mechanical forces at the interface, for example strain at an epitaxial interface,
- (v) weak interactions like van der Waals forces at the interface,
- (vi) interactions between magnetic spins.

Figure 1 shows a schematic for classifying composites, according to the interaction at their interfaces. For example, if only specific molecules can transfer through the interface, then combining this with other functionality, such as catalysis and photocatalysis, could realize molecular selective (photo)catalysts [Fig. 1(a), case (i) in the above classification]. Alternatively, electrons and holes could transfer through the interface between

Fig. 1. Schematic illustrations of functional interfaces: (a) molecular selective (photo)catalysts, (b) electron transfer between semiconductor particles in Z-scheme photocatalysts, (c) two-dimensional electrons induced on an epitaxial interface, (d) electron carriers induced by a strong electrical field on a semiconductor surface, (e) formation of a sponge crystal by epitaxial self-assembly of nanocrystallites, (f) epitaxial film subjected to strain from a single crystal substrate, (g) uniaxial growth of a layered compound on a smooth substrate surface.
particles [Fig. 1(b), case (ii)], or the interface between a thin film and substrate. Photocatalysts that operate by the so-called “Z-scheme” fall into the category shown in Fig. 1(b). When two kinds of semiconductor particles are in contact with each other, one semiconductor may absorb photons to generate excited electrons in the conduction band, which then transfer to the valence band of the other semiconductor, through the interface between the particles [case (ii)]. Electrons induced at the interface of an insulating substrate and epitaxial thin film fall into the category in Fig. 1(c) [case (iii)]. A strong electric field at the interface can induce charge carriers at the semiconductor surface, as shown in Fig. 1(d). The electrical double layer at the interface of a solid and ionic liquid produces an electric field. This field can be sufficiently strong to bring about the transition from a semiconducting surface to a metallic conduction [case (iii)].

Epitaxy between inorganic crystals is another important phenomenon. Nanocrystallites can aggregate in the same crystal direction, with their connection arising homoeptaxially by local dissolution–precipitation. This yields a “sponge crystal” [Fig. 1(e)]. As we discovered in 1996, and we named this process “epitaxial self-assembly”. Hetero-epitaxial growth of thin films on crystalline substrates can also significantly affect the structure and physical properties of the crystalline thin film. For example, the metal–insulator transition temperature of VO2 epitaxial thin films can reportedly be controlled by the tensile or compression strain of the substrate [Fig. 1(f), case (vi)]. Alternatively, if the interaction between the film and substrate is weak, the thin film orientation can be dictated by the nature of the crystalline phase. The basal plane of compounds with layered structures tends to adopt the layered orientation. When the material is inorganic, the layer [Fig. 1(g), case (v)]. This has been exploited to prepare uniaxially oriented thin films of the layered compound Sr2N. This compound is isostructural with Ca2N; a two-dimensional electride compound that has attracted much recent attention. Through the epitaxial interface, magnetic interaction brings about “exchange bias”, in which directions of magnetization of ferromagnetic thin films are controlled by the interaction with the spins at the surface of antiferromagnetic substrate [case (vii)]. Another area of thin film synthesis is in preparing solid solutions. This involves two or more components being supplied in atomic form to the substrate, to achieve mixing of the components at the atomic level.

In this article, focusing mainly on our studies, we provide an overview of the role of interfaces in composites, to achieve various functionality, and to understand the effects of interfaces on the structure and properties of solids.

2. Molecular selective photocatalysts realized by composites with nanoscale structures

TiO2 is a versatile photocatalyst. It can potentially be applied in the production of hydrogen from water, and in the removal of organic contaminants from air and water. In environmental purification applications, TiO2 absorbs ultraviolet wavelengths to generate active species such as hydroxyl radicals and O2-. These active species react with organic contaminants, decomposing them to CO2 and H2O. TiO2 as a photocatalyst has no molecular selectivity, so non-selectively decomposes all organic molecules. This results from the non-selective nature of the active species generated at the surface of the photocatalyst. This can be beneficial, since all organic contaminants can be removed by a single photocatalyst. However, the non-selectivity of the TiO2 photocatalyst can be detrimental when concentrated co-existing species suppress the reaction of a dilute target contaminant. Physiologically active molecules often retain their activity at very low concentrations. A high selectivity toward these molecules is necessary to achieve their complete decomposition by photocatalysis. Molecular selective photocatalysis was first proposed around 2000. Studies of this including our own works soon followed, and were built on by many subsequent studies in the following decade. A review article on this field was published in 2013.

2.1 Combining molecular selective adsorption and photocatalysis

A simple and effective strategy for realizing molecular selective photocatalysis is to combine molecular selective adsorption and photocatalysis. An early photocatalyst model is shown in Fig. 2. Its structure is formed on the surface of a flat substrate. Domains affording molecular selective adsorption sites and domains that are photocatalytically active are located to allow molecules to transfer from the former to the latter.

2.2 Octyl group-grafted TiO2 (C8-TiO2) and octyl group-grafted TiO2/mesoporous silica (C8-TiO2-MPS)

A high surface area promotes high photocatalytic activity, so small crystals and porous materials are advantageous as components of photocatalysts. We exploited a simple strategy to realize molecular selective photocatalysis. Organic molecules were grafted directly on the surface of photocatalytically active particles. This strategy imparted molecular selectivity on the photocatalyst surface. The target molecule was nonylphenol; an endocrine disrupting chemical that is active for fish. Nonylphenol is amphiphilic. It has a large branched nonyl group as the hydrophobic part, and a phenolic hydroxyl group as the hydrophilic part.

Commercially available P25 TiO2 was used as the photocatalyst, as it is well known to be highly active. Figure 3(a) shows a TiO2 surface was functionalized with hydrophobic octyl groups (C8-TiO2). Photocatalytic activity was tested using an aqueous solution containing 2 ppm nonylphenol and 1000 ppm phenol. The solution contained a coexisting molecule (phenol) with a concentration of 500 times higher than the target molecule (nonylphenol). Figure 3(b) shows the time course of the nonylphenol concentration. When the photocatalyst was added to the solution, the nonylphenol concentration noticeably decreased by adsorption. Photoirradiation was then commenced, upon which the nonylphenol concentration decreased rapidly. Nonylphenol was completely removed after several hours, in the presence of concentrated coexisting phenol.

For comparison, P25 TiO2 without surface modification was also tested under the same conditions. The results are shown inset in Fig. 3(b). The decomposition rate of nonylphenol decreased significantly after 1 h of photoirradiation, and significant nonylphenol remained after prolonged photoirradiation, because of the presence of concentrated phenol. This molecular selective photocatalyst exhibited very high activity, because the TiO2 particles were highly crystalline and had high photocatalytic activity.
Despite this high activity and molecular selectivity, the durability of the photocatalyst was poor. The photocatalytic reaction gradually decomposed the surface functionalization, and molecular selectivity deteriorated with prolonged photoirradiation.

We synthesized another nanostructured composite photocatalyst, in which hydrophobic organic groups were grafted inside the pore walls of TiO₂-supported mesoporous silica (C₈-TiO₂-MPS). The molecular selectivity of this photocatalyst was high, but its photocatalytic activity was low, because of low TiO₂ crystallinity and loading (approximately 13 wt. % maximum). The results of these studies indicate that (i) combining molecular selective adsorption and photocatalysis is very effective, (ii) pre-formed highly crystalline TiO₂ particles are preferable as the photocatalytic moiety, and (iii) the photocatalyst should contain only inorganic materials, to maximize catalyst stability.

2.3 TiO₂ particle-mesoporous silica composites

This strategy assigns the roles of molecular selective adsorption and photocatalytic activity to mesoporous silica and pre-formed well-crystalline TiO₂ particles, respectively. The TiO₂ particle size was approximately 20–30 nm, which was much larger than the pore diameter of mesoporous silica of approximately 3 nm. Thus, the TiO₂ particles could not be loaded into the pores. An alternative structure was needed to combine these two functionalities. This was achieved in a composite consisting of TiO₂ particles surrounded by mesoporous silica, as shown in Fig. 4(a). This structure was synthesized by precipitating mesoporous silica using the sol-gel method, in a solution containing dispersed TiO₂ particles. Transmission electron microscopy (TEM) images of the composite are shown in Fig. 4(b). The TiO₂ content in this composite was as high as 60 wt. %.

Results of molecular selective photocatalytic tests are shown in Fig. 4(c). An aqueous solution containing four alkylphenols was subjected to photocatalytic reaction. After adding the composite photocatalyst to the solution, decreases in concentrations due to selective adsorption were observed for nonylphenol and heptylphenol, because of their larger alkyl groups. After photoirradiation, the preferentially adsorbed nonylphenol and heptylphenol were decomposed much faster than the other two phenols. Another experiment demonstrated the importance of molecular selectivity: in the presence of phenol with a concentration 100 times larger than those of nonylphenol and heptylphenol, the two target compounds were completely decomposed by the molecular selective photocatalyst, but were unable to be so when using pristine TiO₂ as the photocatalyst.

These reports stimulated further studies on molecular selective photocatalysis, and a review on this topic was recently published.

3. Nanoscale organic–inorganic hybrid catalysts based on mesoporous silica

3.1 Solid acid catalysts operating in water

Mesoporous silica has a characteristic nanoscale structure, with uniform ordered pores of diameter of several nanometers. Many studies have exploited this structure to realize various functionalities. Mesoporous silica has a high surface area, and many studies have explored its applications in catalysis. The inner-pore walls of mesoporous silica are often functionalized by chemical modification. The following highly active solid acid catalyst has a nanostructure based on mesoporous silica. Organic groups and inorganic molecules were grafted on the inner-pore walls.

Solid acid catalysts have benefits compared to liquid acids such as H₂SO₄, because solid catalysts are easily separated from solution, and are less corrosive to metallic reactors. In aqueous solutions, only a limited number of solid acid catalysts are workable, because acidic protons on solid surfaces are deactivated in water. In 1981, a hydrophobic zeolite catalyst (H-ZSM-5) was reported to be active in aqueous solution. Another solid acid catalyst active in aqueous solution is the acidic inorganic salt Cs₂₅H₄₅PW₁₅O₄₆. It was reported in 1997 that this catalyst exhibited the highest reaction rate per acidic proton at the...
Before 1996, studies had been reported about metal oxides located in hydrophobic reaction fields. However, if acidic protons are completely buried in a hydrophobic environment, water and reactant molecules cannot access the active sites. Grafting long hydrophobic chains on pore walls inhibits liquid water from intruding into pore spaces of mesoporous silica.23) Thus, the nanostructure in Fig. 5 was designed.24) Spaces were left at the centers of pores, to allow free diffusion of the solvent and reactant molecules within pores. Active acidic protons were required to be exposed to the interface between the aqueous solvent and hydrophobic layer that was grafted on the inner pore wall. Thus, octyl groups and H3PW12O40 heteropolyacid inorganic molecules were used as the hydrophobic groups and catalytically active acidic inorganic species, respectively. The size of the inorganic molecule was approximately 1 nm, and the octyl group length was comparable. Thus, a hydrophobic reaction environment formed around the inorganic molecules, and acidic protons on the catalysts exhibited the highest reaction rate per acidic proton. The activity per acidic proton was up to six times higher than that of sulfuric acid, a commercially used liquid acid. The concept proposed in this study, namely of combining a hydrophobic reaction field and inorganic acidic molecules on the nanoscale, was used in subsequent studies of acid catalysts in aqueous media.25,26)

3.2 Nanoscale design strategy

Previous studies hinted at the potential of acidic protons located in hydrophobic reaction fields. However, if acidic protons are completely buried in a hydrophobic environment, water and reactant molecules cannot access the active sites. Grafting long hydrophobic chains on pore walls inhibits liquid water from intruding into pore spaces of mesoporous silica.23) Thus, the nanostructure in Fig. 5 was designed.24) Spaces were left at the centers of pores, to allow free diffusion of the solvent and reactant molecules within pores. Active acidic protons were required to be exposed to the interface between the aqueous solvent and hydrophobic layer that was grafted on the inner pore wall. Thus, octyl groups and H3PW12O40 heteropolyacid inorganic molecules were used as the hydrophobic groups and catalytically active acidic inorganic species, respectively. The size of the inorganic molecule was approximately 1 nm, and the octyl group length was comparable. Thus, a hydrophobic reaction environment formed around the inorganic molecules, and acidic protons on the inorganic molecules were exposed to the interface of the aqueous phase intruding into pores, as shown in Fig. 5. Aminopropyl groups were grafted, to anchor inorganic molecules on pore walls. One inorganic molecule was anchored through one ammonium group formed by reaction with one acidic proton. Therefore, two protons per inorganic molecule remained unreacted, which was confirmed by the adsorption of basic pyridine molecules on the catalyst.

3.3 Catalytic performance

Table 1 summarizes the results of the catalytic reactions. The composite catalyst exhibited the highest reaction rate per acidic proton. The activity per acidic proton was up to six times higher than that of sulfuric acid, a commercially used liquid acid. The concept proposed in this study, namely of combining a hydrophobic reaction field and inorganic acidic molecules on the nanoscale, was used in subsequent studies of acid catalysts in aqueous media.23,24)

4. Sponge crystals: a new class of porous crystals

Before 1996, studies had been reported about metal oxides particles aggregates having characteristic shapes. For example, Sugimoto et al. reported that iron oxide formed pseudocube-shaped aggregates of crystallites.27) The constituent crystallites had the same crystal orientation in an aggregate, but no epitaxial connections formed between the crystallites, unlike the sponge crystals as is described below. At the time there was no report that epitaxial connections forms between crystallites in an aggregate and the whole aggregate is coherent in terms of X-ray diffraction.

In 1996, we reported the ordered structure of porous crystal of (NH4)3PW12O40.6) It exhibited high porosity, but its crystal structure contained no interstitial spaces or built-in pores, such as occur within zeolites.29,30) In 2006, we reviewed our studies of this compound, and named this class of porous crystals, sponge crystals.31) The definition of a sponge crystal in 2006 was: ‘‘Sponge crystals’’ are single crystals containing continuous voids, and unlike zeolites, have no intrinsic structural pores. To comply with this definition of ‘‘sponge crystals’’, the crystal structure of a compound should not have intrinsic pores defined by the crystal structure itself, therefore ruling out zeolites as sponge crystals.’

(NH4)3PW12O40 crystals precipitated at elevated temperatures comply with this definition. Figure 6 shows an SEM image of (NH4)3PW12O40 sponge crystals. Each dodecahedral crystal was highly microporous. The formation process is shown in Fig. 1(e). The ordered porous crystals form by the self-assembly of nanocrystallites. Epitaxial connections form between them by dissolution–precipitation at interfaces, leaving spaces between crystallites as pores. X-ray diffraction and various microscopy techniques supported the characterization of this structure and its formation mechanism. In 1996, we named this crystallite self-assembly mechanism ‘‘epitaxial self-assembly’’.6) Sponge crystals are coherent in terms of X-ray diffraction, because the crystallites are epitaxially connected with each other. (After defining the term ‘‘epitaxial self-assembly’’ in 1996, the same term began to be used with a different meaning in 2003,31) which has caused some confusion.)

After our report of epitaxial self-assembly, Colfen et al. proposed the concept of ‘‘mesocrystals’’.32) This is a wider category than sponge crystals. Mesocrystals have a characteristic shape, which reflects the shape of their constituent crystallites. Thus,
As shown in Fig. 1(g), when the interaction between thin films and substrates is weak, such as in layered compounds, thin films tend to grow with a uniaxial orientation. The basal plane of the layered structure tends to become parallel to the substrate surface. This phenomenon could be exploited to prepare uniaxially oriented thin films of the layered nitride SrN on MgO(001) substrate, as shown in Fig. 8.

The compound SrN has been known for a long time.\textsuperscript{35,36} As shown in Fig. 8, SrN crystallizes in an anti-CdCl\textsubscript{2} structure. It is isostructural to CaN, which has attracted much attention as a two-dimensional electrode compound.\textsuperscript{9} SrN is also a two-dimensional electrode. In 2004, we measured the electronic conductivity of SrN, in the direction parallel to the layers of the compound.\textsuperscript{9} The resistivity was as low as that of Sr metal, and its temperature dependence was metallic. This supported the concept of two-dimensional free electrons, which was proposed for CaN and SrN.\textsuperscript{9,37}

### 5.2 Effects of epitaxial interfaces on substrates

As discussed in the introduction, exploiting the strain from substrates is a promising strategy for controlling the physical properties of thin films. The physical properties are tunable, from those of the bulk crystal. For instance, it has been reported that the metal-insulator transition temperature of VO\textsubscript{2} thin films can be tuned by the strain of the substrate.\textsuperscript{7} We also reported that the growth orientation of CrN epitaxial thin films on MgO(001) and \(\alpha\)-Al\textsubscript{2}O\textsubscript{3}(0001) substrates affected the antiferromagnetic transition of CrN. This was possibly due to strain and/or constraint (i.e. the effect of substrates fixing the positions of atoms at their interfaces, preventing them from changing their interatomic distances) from the substrate.\textsuperscript{38} These examples demonstrate that transitions are accompanied by a change in structural distortion or symmetry.

### 5.3 Solid solution thin films

Thin film synthesis is useful for preparing solid solutions, because two or more components can be supplied in atomic form to the substrate, to achieve mixing of the components in atomic level. We investigated metal nitride solid solutions, including Ti\textsubscript{1-x}Ni\textsubscript{x}N\textsuperscript{39} and Cr\textsubscript{1-x}Ti\textsubscript{x}N.\textsuperscript{40,41} The latter exhibited ferromagnetism and characteristic magnetoresistance behavior. The mechanism was discussed by means of band calculations by a theoreticist.\textsuperscript{42} The behavior of the nitride was compared with that of the colossal magnetoresistance (CMR) of the oxide La\textsubscript{1-x}Sr\textsubscript{x}MnO\textsubscript{3}.\textsuperscript{40}

### 6. Conclusions

The effects of interfaces were overviewed in various inorganic materials and composites, largely consisting of our studies over the last two decades. The materials and phenomena discussed range from porous materials as (photo)catalysts, to the self-
assembly of nanocrystallites, to epitaxial thin films on single crystal substrates. The important point is to control the structure of an interface at the atomic or molecular level, even though characterization of the interface structure is often difficult. Designing functionality into interfaces is a promising strategy for exploring new phenomenon in solid state materials and nanostructured composites.

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References

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