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Selectivity in Ligand Functionalization of Photocatalytic Metal Oxide Nanoparticles for Phase Transfer and Self-Assembly Applications

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Abstract: Functionalization of photocatalytic metal oxide nanoparticles of TiO₂, ZnO, WO₃ and CuO with amine-terminated (oleylamine) and thiol-terminated (1-dodecanethiol) alkyl chained ligands was studied under ambient conditions. A high selectivity was observed in the binding specificity of a ligand towards nanoparticles of these different oxides. It was observed that oleylamine binds stably to only TiO₂ and WO₃, while 1-dodecanethiol binds stably only to ZnO and CuO. Similarly, polar to non-polar solvent phase transfer of TiO₂ and WO₃ nanoparticles could be achieved by using oleylamine, but not by 1dodecanethiol, while the contrary holds for ZnO and CuO. The surface chemistry of ligand functionalized nanoparticles was probed by ATR-FTIR spectroscopy, that enabled to elucidate the occupation of the ligands at the active sites. The photo-stability of the ligands on the nanoparticle surface was determined by the photocatalytic self-cleaning properties of the material. While TiO₂ and WO₃ degrade the ligands within 24 hours under both UV and visible light, ligands on ZnO and CuO remain unaffected. The gathered insights are also highly relevant from an application point of view. As an example, since the ligand functionalized nanoparticles are hydrophobic in nature, they can thus be self-assembled at the air-water interface, for

obtaining nanoparticle films with demonstrated photocatalytic as well as anti-fogging properties.

Introduction

Nanoparticles of semiconductor oxides such as TiO₂, ZnO, WO₃ and CuO have gained considerable attention over the years for their proven usefulness in important applications such as photocatalysis, water splitting, CO₂ reduction, solar cells, super-capacitors, sensing applications, amongst others.^[1-3] Many of these applications require functionalization of the nanoparticles with organic ligands with desired functionalities.^[4] For instance, bio-functionalization of nanoparticles is essential for in-vitro/in-vivo medical applications in order to render them bio-compatible, target-specific^[5] and stable.^[6] Other applications of the functionalization of nanoparticles include molecular imaging^[7], click-chemistry mediated conjugation^[8], selective self-assembly of nanoparticles^[9], ligand accelerated catalysis,^[10] and many more. For nanoparticles synthesized in an aqueous medium, hydrophobic functionalization is often necessary to stably disperse the nanoparticles in a non-polar medium. Therefore, it is important to have

hydrophobic functionalization strategies available that enable phase transfer of nanoparticles from an aqueous to non-polar organic phase, for follow-up reactions in this non-polar phase^[11], coating of hydrophobic substrates^[12], self-assembly as thin films^[13], and so on.

The efficacy of the functionalization is dependent on the affinity of functional head groups towards the nanoparticles' surface.^[4] Chemisorption by the formation of a covalent-like bond results in stable functionalization, while electrostatic or hydrophobic interactions result in a weaker attachment for which the surface coverage is subject to the equilibrium concentration of the ligand in the solution. The exact nature of these bonds is still an important topic of investigation even for an extensively explored case of Au and thiol.^[14,15] Recently for instance, Inkpen *et al.* challenged the widely accepted view that the Au-thiol linkage is covalent showing that the nature of the bond is mainly physisorption.^[16]

Nonetheless, the strong bonding of thiols to both metal and metal-oxide nanoparticles is overall well known. Similarly, amines also bind strongly with metal as well as metal oxide nanoparticles. In heterogeneous catalysis, thiols and amines have therefore been the most explored anchoring ligands. The relative affinity of different functional groups to bind to the nanoparticles is another important aspect as it determines the success of a ligand exchange process to replace any existing ligand on the nanoparticle surface with a new ligand. In this case, the new ligands must have a stronger affinity to bind in comparison to the existing surface ligand.^[17,18] Due to the stronger affinity, oleylamine capping on Au nanoparticles can be effectively replaced by thiol ligands by simply introducing the thiol ligands to the nanoparticle colloid in a thiol-for-amine ligand exchange process.^[19] Similarly, thiol-for-phosphine ligand exchange has also been achieved at room temperature due to such relative binding affinity.^[20]

While functionalization of metal nanoparticles is wellexplored, metal-oxide nanoparticles are far less studied in this context. In this work, we report on functionalization of semiconductor oxide nanoparticles, namely TiO₂, ZnO, WO₃ and CuO with amine (oleylamine) and thiol (1-dodecanethiol) ligands in identical chemical environments. This way the selectivity of a functional group towards a specific oxide becomes clear and is further consolidated by the fact that the efficacy of phase transfer of these nanoparticles from aqueous to non-polar (hexane or chloroform) phase is in direct correspondence with the ability of the ligand *i.e.*, the head group to chemically attach on to the nanoparticles' surface. This work probes the ligand-capped nanoparticles' surface chemistry by (ATR-) FTIR with respect to dissociative and molecularly adsorbed H₂O on active sites, in order to understand to what extent the different ligands occupy the active sites on the various metal oxide surfaces. Importantly, the selection of oleylamine and 1-dodecanethiol for these experiments is based on the fact that the optimum chain length is dependent on the functional group.^[21,22] The alkyl chain lengths, 18-C and 12-C of oleylamine and 1-dodecanethiol, respectively, are considered optimal for phase transfer experiments. Soliwoda *et al.* showed that for thiols, 1octadecanethiol with 18-C alkyl chains form disordered assemblies on the nanoparticle surface which results in poor phase transfer.^[22] While, for amines, it takes longer 18-C chains to facilitate stable and efficient phase transfer of nanoparticles.^[21] In the context of hydrophobization of nanoparticles by selective ligand functionalization, phase-transfer and air-water interfacial self-assembly, the findings from this study provide fresh insights for applications in many of these areas, of which several will be demonstrated.

Results and discussion

Functionalization and phase-transfer of nanoparticles:

The FTIR spectra in Figure 1 compare the presence of the two organic ligands on the different nanoparticle powders after surface adsorption and repeated washing. The intensity of the general v_s (-CH₂-), v_{as} (-CH₂-), and v_{as} (-CH₃) stretches of the alkyl chain in the 2850-3000 cm⁻¹ region indicates the amount of both oleylamine and 1-dodecanethiol ligand present in the

dry powder. The weaker bands at 1466 and 1377 cm⁻¹ arising from the asymmetric and symmetric bending vibration of the methyl group, $\delta_{as}(-CH_3)$ and $\delta_s(-CH_3)$, respectively, also represent the ligands. Repeated washing steps ensure that any weak physisorbed species will be removed due to mass action upon introduction of fresh solvent. Clearly, oleylamine remains bound to TiO₂ and WO₃, while 1dodecanethiol remains on ZnO and CuO nanoparticles. It is, however, difficult to comment on the true chemical nature of the bond on the basis of the stability through washing steps. In the transmission mode, due to the higher opacity of WO₃, WO₃ sample amount had to be kept minimum yet sufficient as the spectra become too noisy for sample amounts same as the other materials.^[23] The nanoparticle-ligand interaction becomes clear in the discussion of phase transfer

experiments and hydrophobicity by water contact angle measurement later in this study. The broad absorption band in the 3200-3800 cm⁻¹ region corresponds to different bands of H₂O adsorbed both dissociatively and in molecular form.^[24] This broad absorption band is significantly weakened by adsorption of 1-dodecanethiol on ZnO nanoparticles. However, absorption of oleylamine does not suppress this absorption band in TiO₂ and WO₃ nanoparticles. Since water is adsorbed in many forms, it is important to differentiate those in order to derive further information, as discussed in section 3.2. Usually, the bands for amine and thiols are difficult to detect due to their relative weakness. ^[25,26]

Both thiol-metal and amine-metal bonding have been known to be strong interactions, while metal oxidethiol or metal oxide-amine bonding has not been



Figure 1. FTIR (transmission) spectra comparing $v_s(-CH_2-)$, $v_{as}(-CH_2-)$, and $v_{as}(-CH_3)$ stretches of the alkyl chain of ligand functionalized nanoparticles after four washing cycles: (a) oleylamine and oleylamine + nanoparticles (b) 1-dodecanethiol and 1-dodecanethiol + nanoparticles.

explored extensively.^[27,28] In ligand-exchange studies, thiol often successfully replaces other functional groups under ambient conditions.^[18,4] For instance, it has also been shown that octadecanethiol can effectively replace oleylamine on Au nanoparticle surface by ligand-exchange.^[29] Generally, reports on controlled experiments comparing binding tendencies of thiols and amines on metal or metal oxide surfaces are scant.^[30] However, nanoparticle considering the known strong dependency of ligand interaction on the crystal planes^[31], lewis- Brønsted acid sites^[32], etc., the selectivity of both oleylamine and 1-dodecanethiol to these four metal-oxides in Figure 1 is an expected occurrence. It has been hypothesized that ligands can coordinate as an anion to excess metal atoms on the surface to balance charge and terminate the lattice (X-type), such as oleic acid, dodecanethiol, or phosphonic acids, or as a neutral dative bond (Ltype, Lewis basic), as in oleylamine, trioctylphosphine oxide, and trioctylphosphine.^[33] To explain the reciprocity in the trend shown in this study, deeper insight into the atomistic interaction is required to explain the tendency of a certain functional group to a certain surface, which is a subject of ongoing theoretical/computational investigations.^[34]

This selectivity of one type of functional group to a certain metal oxide observed in the adsorption experiments explains success of the phase transfer experiments of nanoparticles with a specific ligand only, as illustrated in Figure 2, and with the results of the different metal oxide-ligand phase transfer experiments summarized in Table 1. Very clearly, phase transfer with oleylamine only works for TiO₂ and WO₃, while 1-dodecanethiol only works for ZnO and CuO. Figure S1 shows the inability of the non-attaching ligands to successfully transfer nanoparticles from a water/ethanol phase to a hexane phase. The phase transfer process requires strong nanoparticle-ligand bonding in order for the nanoparticles to cross the interfacial barrier between the polar and the non-polar phases.^[35,36] Apparently, the amine-ZnO or -CuO and thiol-TiO₂ or -WO₃ interactions are too weak to permanently hydrophobize the nanoparticle surfaces for the interfacial transfer. The thiol-ZnO or CuO bond

are strong, resulting in smooth phase transfer and very pronounced infrared absorption bands even after repeated washing steps. It has been shown with XPS analysis that both Zn- and O-terminated nanoparticles of ZnO bind quite strongly with thiol groups and these bonds are thermally stable up to 350 °C.^[37] Interestingly, on ZnO nanoparticles, an even stronger affinity of phosphonic acid functional groups as compared to thiols has been reported.^[38] While certain reports on amine functionalization of ZnO and CuO nanoparticles can also be found in literature^[39,40], it is clear from our experiments that the interaction is not strong enough for phase transfer. On the other hand, the strong interaction between oleylamine and TiO₂ or WO3 easily enables phase transfer. It is important to note that a ligand that binds strongly to TiO₂ and WO₃ does not necessarily have to be an ineffective binder to ZnO or CuO surface. It is known that organic ligands with phosphonic acid functional head groups bind strongly to both TiO₂ and ZnO.^[38,41,42,36] This consolidates the fact that the nature of the ligandnanoparticle interaction varies case by case and the selectivity is determined by both the ligand and the nanoparticle surface chemistry. It is, however, also important to have a sufficiently high ligand concentration as the contact time needed for effective attachment is inversely proportional to this concentration.^[43,44]

Table 1. Summary of phase transfer experiments ofeach metal-oxide nanoparticle with oleylamine and 1-dodecanethiol.

	oleylamine 1-dodecanethiol	
TiO ₂	successful	unsuccessful
WO ₃	successful	unsuccessful
ZnO	unsuccessful	successful
CuO	unsuccessful	successful

The selectivity of ligands to bind a given oxide also prompts experimentation to selectively transfer a



Figure 2. (a) Schematic explanation of the polar to non-polar phase transfer procedure of nanoparticles (inset: HAADF-STEM images of TiO_2 nanoparticles before and after phase transfer with 100 nm scale bar). (b) Successful phase transfer of each nanoparticle with its specific ligand.

particular oxide nanoparticle from a mixture of two. The trial experiment in Figure S2 shows that 1-dodecanethiol can selectively transfer CuO and ZnO nanoparticles from a mixture of CuO-TiO₂ and ZnO-TiO₂, respectively. However, selective phase transfer of TiO₂ using oleylamine was not possible possibly due to the lack of a relative binding tendency as strong as in the case of thiol. It indicates that thiol-binding is more chemically selective.

Surface chemistry of nanoparticle powders

Since adsorption of H₂O on metal oxide surfaces, especially TiO₂ and ZnO, is well known, comparison of pure nanoparticle powders with ligand functionalized powders with FTIR provides important insight into the ligand binding on the surface. Since the transmission mode includes the H₂O peaks possibly due to capillary condensation in the pores, the attenuated total reflectance (ATR) mode is more appropriate for such

surface chemistry characterization. In general, ATR-FTIR is a rather simple yet efficient method for surface characterization, especially in the case of WO₃ that yields poor spectral information in the transmission mode due to its opacity. The broad absorption band in the FTIR spectrum of both pure TiO₂ and ZnO powders in the region 3000-3700 cm⁻¹ is commonly assigned to the stretching vibration modes of H₂O molecules, Figure 3(a).^[45,46] In the context of surface chemistry, the band at 3695 cm⁻¹ is of particular interest as it represents isolated non-hydrogen-bonded OH groups on TiO₂, as shown in the HREELS study by Henderson, indicating availability or unavailability of active sites.^[47] interaction per H₂O molecule is not strong.^[48] Lu *et al.* attribute the absence of the H₂O peaks on commercial WO₃ to low surface area.^[49] Similarly, a number of DFT studies has shown the possibility of both molecular and dissociative adsorption of H₂O on CuO surfaces^[50,51], in contrast to the FTIR spectra in Figure 3, which can in fact be attributed to low surface area. Although surface adsorbed water on WO₃ and CuO is not apparent from the ATR-FTIR spectra, the presence of trapped moisture in the pores has been indicated in the transmission spectra in Figure 1 and TGA analysis in Figure 4. Nevertheless, comparison with respect t the surface adsorbed H₂O is possible only in the cases of TiO₂ and ZnO.



Figure 3. ATR-FTIR spectra of ligand functionalized nanoparticles (red) compared with their pure powders (black) in the (a) 3800-3100 cm⁻¹ (b) 2500-1000 cm⁻¹ regions in terms of H_2O adsorption bands.

known to be from the scissoring modes of molecularly adsorbed H_2O , Figure 3(b). In contrast, the surfaces of WO₃ and CuO nanoparticles are free from adsorbed water. It has been shown by Albanese *et al.* with the help of DFT calculations that H_2O preferably adsorbs on WO₃ in its un-dissociated form due to the presence of ions at the surface that act as Lewis acid sites and this Since H_2O adsorption on TiO_2 and ZnO has been studied extensively, it is useful to discuss these cases with respect to the ATR-FTIR results in order to understand the ligand-nanoparticle interaction. On the (101) surface of anatase TiO_2 , which is dominating in this form, molecular adsorption of H_2O is energetically favorable in comparison to dissociative adsorption as

shown by Vittadini et al.^[52] This stability is thought to be due to the hydrogen bonding between the H atoms of the adsorbed H₂O molecules with bridging O atoms on TiO₂. On the (001) surface, dissociative adsorption is favored up to half coverage of the Ti sites beyond which molecular adsorption is more likely.^[52] Physical adsorption of H₂O on the first layer via hydrogen bonding has also been shown to be an energetically favorable occurrence. Similarly, on the energetically favorable $(10\overline{1}0)$ surface of ZnO, H₂O adsorbs both dissociatively and as molecules.^[53,46] In the 3100-3800 cm⁻¹ region of broad band H₂O absorption in Figure 3(a), as discussed already, the shoulder at 3695 cm⁻¹ in both pure TiO₂ and ZnO distinctly signifies dissociatively adsorbed water to the surface.[47,24,54] The absence of this shoulder in the spectra of oleylamine-TiO₂ and 1-dodecanethiol-ZnO nanoparticles indicates unavailability of these sites for dissociative chemisorption of H₂O. Similarly, the bands at 1637 and 1630 cm^{-1} in pure TiO₂ and ZnO, respectively, indicate these molecular adsorbed water bands are broadened or hidden in the ligand capped TiO₂ and ZnO. This also indicates occupation of these adsorption sites by the ligands. The 1466 cm⁻¹ band for the, δ_{as} (-CH₃) asymmetric vibrations in both oleylamine and 1-dodecanethiol is also quite clear on ligand grafted nanoparticles. This band is particularly strong in 1-dodecanethiol-capped ZnO and CuO nanoparticles

in contrast to the weak intensity in oleylamine-capped TiO_2 and WO_3 nanoparticles. As these experiments were conducted in the dark without any chance of photo-degradation, this indicates weaker oleylamine-nanoparticle interaction and reduction of surface-ligands due to mass action of washing steps.

Thermogravimetric analysis was carried out to gain further insight into the ligand-functionalization of the nanoparticles, Figure 4. 1-dodecanethiol starts to burn off at a lower temperature (~150 °C) than oleylamine (200 °C) below which both the compounds remain at their fixed initial weight, Figure S3. For the dry powders, the weight loss due to moisture loss starts at around 50 °C in all the samples. Also, for the oleylamine capped TiO₂ and WO₃, the weight loss already begins at >50 °C, hence the presence of significant trapped moisture despite ligand functionalization is evident. In the transmission FTIR spectra in Figure 1, this trapped water also contributes substantially to the 3000-3500 cm⁻¹ broad absorption band. In contrast, such an early weight loss is not observed in 1-dodecanethiol capped ZnO and CuO nanoparticles, meaning that moisture is present in rather low amount, which is in line with the FTIR spectra of thiol-capped ZnO and CuO nanoparticles where the H₂O band intensity is significantly reduced as compared to uncapped (as oleylamine is not present) ones. This has as a possible



Figure 4. Thermogravimetric analysis (TGA) of (a) oleylamine functionalized nanoparticles (b) 1-dodecanethiol functionalized nanoparticles. Right y-axis in Figure (a): WO_3 powder, right y-axis in Figure (b): CuO + 1- dodecanethiol

implication that 1-dodecanethiol is able to cover ZnO and CuO nanoparticles more stably and extensively. In Figure 4, both oleylamine and 1-dodecanethiol attached on the nanoparticles start burning off at a temperature higher than their pure forms implying strong ligand-nanoparticle bonding. This elevation of the thermal decomposition temperature is again particularly well-defined for 1-dodecanethiol-capped CuO and ZnO nanoparticles due to strong binding. Importantly, for the same ligand functionalization procedure, CuO nanoparticles carry a considerably high weight % (>50%) of 1-dodecanethiol. A possible implication of this may be formation of ligandnanoparticle networks as gel-like consistency was observed for ligand-capped CuO nanoparticles. For TiO₂ and WO₃, the transition from the moisture-loss phase to oleylamine decomposition phase is smooth. It indicates that oleylamine-nanoparticle bonding is not

as strong as that in the case of 1-dodecanethiolnanoparticle. A relatively weaker interaction between TiO_2/WO_3 and oleylamine has already been evident from the FTIR spectra comparisons.

Photo-stability of ligand capped nanoparticles

Given that the semiconductor metal-oxide nanoparticles of TiO_2 , ZnO, WO_3 and CuO are known to show photocatalytic activity, the photo-stability of the ligand-capped nanoparticles under UV and visible light is an important aspect in any possible application. Conversely, as a positive side-effect such high instability, *i.e.* fast photodegradation of the ligands, implies high self-cleaning tendencies of the nanoparticles, which in turn opens up various application opportunities. The photo-stability of the



Figure 5. FTIR spectra (v_s (-CH₂-), v_{as} (-CH₂-), and v_{as} (-CH₃) bands in the 2800-3000 cm⁻¹ region) of ligand capped nanoparticle films on a Si substrate showing photo-stability of the ligands on the nanoparticles under (a) UV (b) visible light.



Figure 6. Degradation curves of oleylamine and 1-dodecanethiol on (a) TiO₂ and (b) ZnO films respectively fitted to a first order kinetic model. The relative concentrations are estimated from the area of the v_s (-CH₂-), v_{as} (-CH₂-), and v_{as} (-CH₃) bands in the 2800-3000 cm⁻¹ region.

ligand capped nanoparticles was tested by monitoring the $v_s(-CH_2-)$, $v_{as}(-CH_2-)$, and $v_{as}(-CH_3)$ bands of the ligands in the 2850-3000 cm⁻¹ region under illumination with UV and visible light, Figure 5. After 24 hours of UV illumination, TiO₂ completely degrades the oleylamine ligands, while WO₃ degrades 45%, with further degradation up to 85% after 48 hours. In contrast, ZnO and CuO nanoparticles could not degrade the ligands to any considerable extent. Similarly, also under visible light (using a white fluorescent tube lamp), both TiO_2 and WO₃ nanoparticles degrade the ligands effectively, while ZnO and CuO are unable to do the same. As expected, the photocatalytic activity of TiO₂ is lower under visible light with 62% degradation in 24 hours, while WO₃ is equally active under both UV and visible light over the 24 h time span (48% degradation in 24 hours). It is important to note that TiO₂ degrades the ligands under UV within an hour (Figure 6); thus, the visible light activity of TiO₂ is negligibly low in comparison to the UV light activity. The inactivity of ZnO, which is also a known photocatalytic material, is counterintuitive. It is reasonable to assume that the strong interaction of thiol ligands results in poisoning of the catalyst surface. Therefore, selectivity

enhancements with thiol ligands are accompanied by a loss of activity.^[55] To understand whether it is the inactivity of ZnO itself or a strong thiol-ZnO binding that prohibits the degradation mechanistically, degradation studies were conducted for both oleylamine and 1-dodecanethiol on TiO₂ and ZnO films on Si wafers, Figure 6. Instead of pre-functionalizing the nanoparticles with the ligands, oleylamine and 1dodecanethiol was spin coated on the surface of both the TiO₂ and ZnO films. Clearly, TiO₂ degrades both oleylamine and 1-dodecanethiol rapidly within an hour. In the case of ZnO nanoparticles, the ligands remain almost unaffected even after 4 hours of illumination. The fitting of the data points to a first order kinetic model shows that the rate constant for ZnO is two orders of magnitude smaller than that for TiO₂. Thus, the 1-dodecanethiol capped ZnO and CuO nanoparticles are photo-stable due to their photocatalytic inactivity. On the other hand, both TiO₂ and WO₃ self-degrade the ligands effectively under both UV and visible light.

Wettability and self-assembly

High wettability, *i.e.* super-hydrophilicity of TiO₂,^[56] WO₃^[57] and CuO^[58] is ZnO. known and hydrophobization of these surfaces is important in many applications. This super-hydrophilicity has mostly been attributed to a photo-induced effect explained by several surface chemistry mechanisms.^[59] For more details on these mechanisms, the reader is encouraged to consult the comprehensive review by Fujishima et al. In addition, the switching between hydrophilic and oleophilic behavior is unique to TiO₂. Photo-induced hydrophilicity (PIH) has also been studied for ZnO, WO₃, and CuO. Like in the case of TiO₂, numerous studies have shown induction of hydrophilicity in ZnO and WO₃ films within a couple of minutes to an hour, while CuO has not been found to show any photo-induced hydrophilicity.^[60-62] On So, the wettability of the ligand-capped nanoparticles of TiO₂, ZnO, WO₃ and CuO is an important aspect as the presence of the hydrophobic alkyl chains alters the surface chemistry. First, the water contact angle on drop-casted films of non-functionalized bare nanoparticle films of all four oxides were found to be close to 0°, Figure 7(a). This may be a photo-induced process as the films were prepared under ambient light. While PIH is not observed in the case of CuO, it also appears highly hydrophilic in nature. The hydrophilicity of all the different nanoparticles is confirmed by contact angle measurements which also explains easy dispersion of these nanoparticles in water to form stable colloids. In contrast, all the ligand capped nanoparticle films are strongly hydrophobic with an average water contact angle >120°, Figure 7(b).



Figure 7. Water contact angle on drop-casted ligand capped nanoparticle films: (a) ligand free bare nanoparticles (b) hydrophobic functionalized nanoparticles, also showing the effect of UV irradiation. The contact angles are average of five measurements and the blue lines marking the angles are for representation.

nanoparticle films such as P25 (TiO₂), where the morphological irregularity may help hydrophobicity, photo-induced super-hydrophilicity can be achieved rapidly, as also verified by our experiments.^[63]

From the experiments in Figures 5 and 6, the irradiation of UV light is supposed to degrade the ligands especially on TiO_2 and WO_3 to revert the surfaces back to their hydrophilic ground state, at a

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rate corresponding to the photocatalytic self-cleaning rate. Thus, the super-hydrophilicity of TiO_2 film after 3 hours of irradiation is consistent with our photostability experiments showing that the ligands are completely degraded under UV within that time period. Similarly, since 24 hours of UV treatment only partially decomposed the ligands on WO₃ the samples in dark for 24 hours, Figure S4. Importantly, while the correlation holds in this particular case, in general photocatalytic activity cannot always be directly related to photo-induced hydrophilicity as it has been shown that despite comparable photocatalytic activity of SrTiO₃, photoinduced hydrophilicity was not observed like in the



Figure 8. Images of assembly of ZnO (a) and TiO₂ (b) nanoparticles at the air-water interface. (c, d, e, f) are SEM images of self-assembled films of TiO₂, WO₃, ZnO (sample 1) and ZnO (sample 2). Sample 1: film after incomplete surface coverage; sample 2: film after complete surface coverage.

nanoparticles (Figure 5a)), the water contact angle is still 86° after 20 hours of illumination. For both ZnO and CuO, the nanoparticle films remain hydrophobic throughout the UV treatment showing consistency with the photo-stability experiments. To confirm the effect of ligand photo-degradation on contact angle, contact angle measurements were taken after keeping

case of TiO₂.^[64]

The contact angle measurement shows that the ligand capping renders the nanoparticles hydrophobic. Thus, self-assembly of these nanoparticles on a water surface is possible by trapping them at the air-water interface which is otherwise not possible, Figure S5. In Figure 8 (a), a partially covered water surface with ZnO



Figure 9. Antifogging property of a self-assembled TiO₂ nanoparticle film immobilized on glass slide.

nanoparticles is shown. As more nanoparticles are introduced, the self-assembled islands rearrange and eventually develop a continuous film. The selfassembled films obtained with ZnO nanoparticles as shown in Figure 8(e) and (f) have high uniformity. Selfassembled films of TiO₂ and WO₃ nanoparticles, Figure 8(c) and (d), however, are not as uniform due to the presence of agglomerates. While there is high surface coverage in all the cases, the uniformity of the selfassembled films is subject to the morphological uniformity of individual nanoparticles. In all these three cases of TiO₂, WO₃ and ZnO, the morphological and dimensional irregularities of the nanoparticles limit the quality of the films in terms of uniformity. Table 2 summarizes the nanoparticle average sizes as provided by the manufacturer data, primary crystallite size and self-assembled film thicknesses. Film thickness profiles were acquired by stylus profilometry (Figure S6), which captures the transition as the stylus slides from the bare substrate to the film. The profilometry results capture the surface morphology only to an extent as the contact between the needle and the film leads to unavoidable displacement of some particles due to the softness of the assembled layer, hence the relatively large experimental error. Still, the measured average layer thicknesses correspond very well to the expected thickness based on a self-assembled monolayer of the corresponding particles. Even with

these irregular nanoparticles, such self-assembled films can be useful in many applications. As an example, Figure 9 demonstrates the use of such a self-assembled TiO_2 nanoparticle film for antifogging applications.

Table 2. Summary of nanoparticle sizes reported by themanufacturer, primary crystallite sizes and self-assembled film thicknesses.

	Avg. Particle	Avg.	Approx. Film	
Material	size (nm),	crystallite	thickness	
	manufacturer	size (nm) ^a	(nm) ^ь	
TiO ₂	≤25 nm	25.8± 0.4	21.2 ± 4.3	
WO ₃	≤50 nm	48.4 ± 7.6	67.9 ± 13.2	
ZnO	≤40 nm	44.3 ± 5.7	59.3 ± 3.1	

^a Calculated from XRD using the Scherrer equation

^b measured by stylus profilometry

Conclusion

Functionalization of TiO₂, ZnO, WO₃, CuO nanoparticles with alkyl-amine (oleylamine) and alkyl-thiol (1dodecanethiol) ligands was studied under ambient conditions. It was observed that oleylamine binds strongly to TiO₂ and WO₃, but not to ZnO and CuO. Conversely, 1-dodecanethiol strongly attaches to ZnO and CuO, but not to TiO₂ and WO₃. Polar to non-polar phase transfer experiments confirm these trends. The ligand attachment is stable through the impact of mass action in the washing steps, and confirmed by FTIR

data. In the case of TiO₂ and ZnO, it could be shown that oleylamine and 1-dodecanethiol effectively occupy the active sites that are otherwise occupied by OH groups in the pristine powders resulting from dissociative adsorption of ambient H₂O. Under UV and visible light, the stability of the surface ligands on the nanoparticles was found to be dependent not only on the intrinsic photocatalytic activity of the material, but also on the ligand bond strength. While TiO₂ degrades oleylamine within two hours and WO₃ takes more than a day, ZnO and CuO appeared to be quite inactive in degrading the thiol-ligands even after 48 hours of illumination of UV or visible light. Similarly, the hydrophobicity attained by the nanoparticles can be removed by photocatalytic degradation of the amine ligands on TiO₂ and WO₃, while 1-dodecanethiol functionalized ZnO and CuO nanoparticles remain stable and hydrophobic under UV and visible light illumination over an extended period of time (>48 hours). The functionalized nanoparticles can be easily self-assembled at the air-water interface to obtain nanoparticle films for various applications, such as antifogging layers as a demonstrated proof-ofprinciple in this work. While the ligand selectivity of different metal oxide nanoparticles and the corresponding phase transfer processes shown in this work is an important aspect relevant to various application scenarios, the demonstration of the use of these hydrophobic functionalized nanoparticles for self-assembled films is potentially valuable to future research on photocatalytic surfaces, photoelectrochemical and photovoltaic applications. Especially in view of the ongoing research on hybrid multifunctional nanoparticles, self-assembly methods are important for obtaining films in a nonintrusive way.

Conflicts of interest

There is no conflict of interest to declare.

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Keywords: ligand functionalization • ligand selectivity • metal oxide nanoparticles • phase transfer • selfassembly

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Entry for the table of contents



The selectivity in ligand functionalization of photocatalytic metal oxide nanoparticles namely, TiO_2 , ZnO, WO_3 and CuO, was demonstrated for thiol (1-dodecanethiol) and amine (oleylamine) terminated ligands. The success of polar to non-polar phase transfer of nanoparticles is consistent with the ligand-nanoparticle selectivity. Infrared spectroscopy provides important insight into the ligand-nanoparticle interaction. The hydrophobic functionalized nanoparticles can be self-assembled at the air-water interface for thin-film applications.

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