

The Role of Stable Bicarbonate Formation on the Loss of Photocatalytic Activity of TiO₂ in Grout Media

Mert Mehmet Oymak¹ 🝺, Deniz Uner* 🖂 🗈

Middle East Technical University Chemical Engineering Department, Ankara, 06531, Turkey,

Abstract: In this study, we report the photocatalytic activity of TiO_2 monitored by benzene oxidation in the grout medium. The results of the batch reaction tests indicated that the activity of TiO_2 coated on grout was substantially less than TiO_2 coated on a glass substrate. CO_2 adsorption on these samples were monitored by DRIFTS. The results reveal that the loss of activity in the grout medium was due to formation of stable carbonates-bicarbonates in highly alkaline grout media.

Keywords: Photocatalytic benzene oxidation, CO₂ Adsorption/DRIFTS, cement/grout media, dimeric form/bicarbonate, HLW/TiO₂

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*Corresponding author. E-mail: <u>uner@metu.edu.tr</u>.

INTRODUCTION

There is a growing market for self-cleaning and photocatalytic air purifying cementitious materials (1). Similar to cement matrix, grout matrix is also an ideal surface for photocatalytic utilization. Compared to cement matrix, grout matrices are frequently used in indoors and therefore sunlight exposure of these materials is low. However, grout applications such as patios, mosaic, stone, and tile works are aesthetic and high cost surfaces, and self-cleaning properties are desirable. Photocatalysts can be applied onto different structural supports (2), embedded in bulk (3-4). Nowadays, commercial building materials are directly coated with photocatalysts (5). The durability of the photocatalytic building materials is of both academic and commercial concern (6).

Concrete matrix, frequently encountered in buildings, is a highly alkaline environment. This high alkalinity can result in extensive amounts of CO_2 and NOx ad(ab)sorption. A recent study (7) reports that CO_2 absorption in cement matrix can compensate the CO_2 footprint of the cementitious manufacture (5). TiO₂ can also be modified with

alkaline structures to increase CO₂ adsorption. Modification of TiO₂ with NH₄OH and KOH was reported to increase the total CO₂ adsorption capacity by a factor of 9 compared to the untreated sample (8). Furthermore, acidic/basic character of cementitious base materials can influence the activity of the photocatalysts. Kozlov et al. studied photocatalytic degradation of benzene and acetone with H₂SO₄ and NaOH treated samples to observe that high alkaline treatment decreased the acidic sites and photocatalytic activity of the samples (9-10). Strini et al.(11) studied photocatalytic oxidation of BTEX (Benzene, toluene, ethyl benzene and oxylene) using P25 in Portland cement samples, comparing the activities between pure TiO₂ and cementitious materials. TiO₂ added Thev observed 3-10 times decrease in photocatalytic activity of P25 in cement samples when compared to pure TiO₂ activity.

Surface carbonates-bicarbonates are formed on TiO_2 surfaces under CO_2 exposure (12-21). It is known that carbon deposition changes the photocatalytic activity on TiO_2 surfaces under UV exposure or dark conditions (22). Strong Lewis acid (Ti^{4+}) and Lewis base (O^{2-}) sites favor the

¹ Present address: Üsküdar University Chemical Engineering Department, İstanbul, 34662, Turkey

formation of bidentate carbonates and bicarbonate species on the surface, whereas monodentative carbonates are favored by Ti^{3+} sites (23). During photocatalytic benzene decomposition, the source of surface carbonates are the byproducts finally leading to CO and CO₂ as gaseous products (24-27).

In this article, we report activity loss due to the interactions between a commercial TiO_2 photocatalyst and its cement based environment. Our work reveals a link between carbonate-bicarbonate formation and photocatalytic activity loss on the alkaline environment of cement based samples.

EXPERIMENTAL/METHODOLOGY

A commercial TiO₂ sample (Anatase Sachtleben Hombitan LW, will be referred to as HLW from this point onwards) with a specific surface area of 11 m^2/g was used in the tests. HLW -grout sample was prepared by mixing with the grout mortar for the in-grout samples, 33 wt% water was added and the final slurry was cast in a plastic vessel (8.5 cm ID, 5 mm depth). A plain grout mortar was also prepared. The samples were cured for 28 days in a controlled atmosphere conditioned at 23 °C and 50% RH. On-the-grout sample was prepared by depositing 0.01 g TiO₂-deionized water solution using a syringe. On-the-glass sample was prepared by doctor blade method. The samples were air-dried for one day.

Photocatalytic benzene oxidation reaction was carried out in a homemade glass manifold (215 ml) operating in batch mode at 1 atm and room temperature. Prior to reaction measurements, the system was evacuated for 30 minutes and the reaction cell was photo-irradiated for 1 hour under vacuum. 0.2 µL of benzene was introduced to a heated manifold kept at 85 °C through a septum injection port and allowed to evaporate. Benzeneair mixture in gaseous form was transferred from heating system to the reaction cell through a vacuum-tight valve. The reaction products were periodically sampled through a septum by a gastight syringe and analyzed using a Gas Chromatograph (Varian 3900) equipped with FID and PoraplotQ capillary column. 100 W UVA (~365 nm) black light was used for photoexcitation. On the same sample, both dark and UV irradiated activities were measured. The dark experiments were done by covering the sample cell with aluminum foil while keeping the cell illuminated, to maintain identical thermal conditions.

DRIFTS (Diffuse Reflectance Infrared Fourier Transform Spectrometry) studies were performed using a Perkin Elmer (Spectrum 100 Series) Spectrometer equipped with a Pike DIFFUSIR[™] DRIFTS cell connected to a home built gas manifold capable of holding vacuum up to 10⁻⁵ Torr connected to a Varian turbo molecular pump station. Equal amounts of samples were used in the DRIFTS cell and they were in powder form for pure TiO₂, and in the precast form for grout containing samples, which is explained in the first paragraph of experimental section. Prior to the measurements, the manifold and the cell were evacuated for 30 minutes. Subsequently, CO2 was dosed onto sample, while monitoring the pressure by a Baratron gauge (MKS). DRIFTS spectrum was recorded after allowing system to equilibrate for 20 min. After adsorption experiments were completed, the sample was evacuated for 10 min and DRIFTS spectra were also recorded under vacuum.

RESULTS AND DISCUSSION

The photocatalytic benzene oxidation rate was measured through monitoring the concentration as a function of time (Fig. 1a). The rate estimations were based on a pseudo first order kinetics, for practical purposes without making any reference to mechanism. The time dependent disappearance of benzene from the batch reaction chamber was approximated as,

$$\ln\left(\frac{C_a}{C_{a,0}}\right) = -kt \tag{1}$$

The reaction rate constants k can be calculated from the slope of trend lines of $\ln\left(\frac{C_a}{C_{a,0}}\right)$ vs.

time graphs (Fig. 1b). The comparison of pseudo first order rate constants provides a common basis for comparing the activities of different photocatalysts and different environments (28-29).

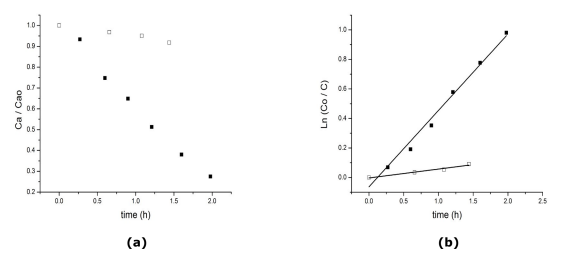


Fig. 1: a) Benzene concentration during photocatalytic oxidation in the grout samples. The filled symbols indicate UV irradiated samples while empty symbols represent the corresponding measurements in dark. 10 wt % TiO_2 was added in the grout in both cases. b) Fig 1a is shown in the

form of $\ln\left(\frac{C_{a,o}}{C_a}\right)$ vs. time.

In order to differentiate the contribution from the cement matrix, similar measurements were performed by coating the TiO_2 samples on the glass. The results were compared with the measurements of the photocatalytic activity of TiO_2 coated on the grout. The results are presented in Table 1. A comparison of the data reported in Table 1 reveals that there is substantial loss of activity on the grout.

According to the findings of an earlier publication (30) from our group, CO_2 evolution was always slower than the disappearance of C_6H_6 , indicating

some carbon hold-up in the structure. To test this hypothesis on the present samples, DRIFTS spectra for various HLW-grout surfaces and pure HLW and grout samples were collected (Fig. 2). IR assignments of adsorbed CO₂ on TiO₂ were made based on the literature as summarized in Table 2. The bands in DRIFTS spectra presented in Figure 2 was assigned as follows: 1800-1200 cm⁻¹ region shows carbonate-bicarbonate related peaks. The absorbance in this region is low for pure grout samples. HLW integration to grout increased adsorption of CO₂ and formation of carbonates-bicarbonates substantially (Fig.2c).

		TiO ₂ surface	kныw (1/h-g cat)				
	density (g/cm ²)						
	On the grout	0.0005	56.6 (±16.0)				
		0.0013	51.3				
		0.0013	196				
	On the glass	0.0020	144				

Table 1: Activity comparison of HLW on glass and on grout surfaces.

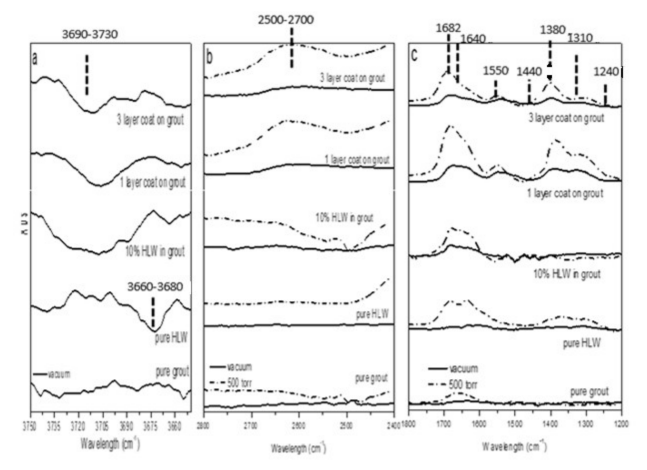


Fig. 2: DRIFTS spectra comparison of pure powder, coating and grout applications using HLW a) 3750-3650, b) 2800-2400, c) 1800-1200 cm⁻¹ regions. The solid lines are collected under 500 Torr of CO₂ while dashed lines indicate the intensity after evacuation. 500 torr CO₂ data for 3750-3650 cm⁻¹ region is not shown due to dominant characteristic CO₂ peaks in the region.

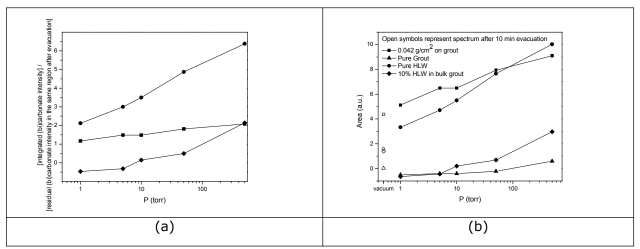


Fig. 3: a) Pressure vs. integrated intensities for 1800-1200 cm-1 region. b) Pressure vs. [Integrated (bi)carbonate intensity] / [residual (bi)carbonate intensity in the same region after evacuation]. Pure grout data in Fig 3b is not shown.

The broad peak around $2500-2700 \text{ cm}^{-1}$ is assigned to dimeric interactions between H and OH groups of bicarbonates (Fig.2b, Table 2). This peak is observed only when HLW and grout were

in intimate contact. This peak was neither observed for pure HLW nor for pure grout sample under CO_2 environment. The decrease in OH population is observed around 3700-3730 cm⁻¹

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for HLW-grout and 3660-3680 cm⁻¹ for pure HLW samples as negative peaks in the regions indicated. This decrease was attributed to the formation of bicarbonates over OH groups upon CO₂ adsorption. The stability of the carbonate species upon evacuation were also monitored. Surface carbonates were found to be more stable on HLW-grout than pure HLW or pure grout samples (Fig.2c). From the data in Figure 2c, it can be seen that the bands in 1200-1400 $\rm cm^{-1}$ range disappear upon evacuation for pure grout and 10 wt% HLW in the grout samples. These bands are however partially stable upon evacuation for pure HLW and HLW on the grout. For a semi-quantitative analysis, the area under the curve for 1800-1200 cm⁻¹ region was integrated and plotted as a function of the pressure (Fig. 3a). It can be clearly seen from the isotherm data in Figure 3a that there is not an appreciable amount of (bi)carbonate hold-up on the pure grout. When $\text{Ti}O_2$ is present, the (bi)carbonate hold-up increases (all the rest of the samples). The relative strength, i.e. stability, of the species was tested against evacuation; open symbols in Figure 3a indicate the residual intensity of the peaks after 10 min of evacuation. In Figure 3b, the isotherm data of Figure 3a (filled symbols) were plotted after they are normalized with respect to their corresponding values under vacuum (open symbols). The same data were also presented in Table 3 in terms of vacuum to 500 torr CO₂ integrated intensity ratios. The normalized values indicate that when HLW and the grout were in intimate contact, these samples hold more surface carbonatebicarbonate species than either pure HLW or pure samples under vacuum conditions. grout However, the data presented in Fig. 3a and Fig. 2c clearly indicates that the bicarbonate formation characteristics on pure HLW and 0.042 g HLW/cm² coating on the grout are similar. The intensity of the bicarbonate species on pure HLW and HLW on the grout, reported in Fig. 3a, is much higher than both the corresponding intensity of pure HLW and that of HLW coated on the grout samples. The differences in the surface

coverages upon evacuation were attributed to the stability of the dimeric bicarbonates.

The DRIFTS results were interpreted as there is a likelihood that a high amount of carbonatebicarbonate species remain on TiO_2 -grout matrix. The photocatalytic benzene oxidation results were interpreted as there is a significant activity loss of TiO_2 in the grout. These two observations are combined broadly to conclude that there is a surface poisoning due to stable carbonate-bicarbonate species in alkaline media.

The formation of bicarbonate species indicate the presence of basic OH groups (14). In this study, dimeric bicarbonate species were formed, which may be a sign of close proximity of OH groups on the surface of HLW-coated-grout matrix samples (Fig. 2b). These species are particularly resistant to evacuation. In addition, for the same samples, OH frequencies shift from 3660-3680 to 3730-3700 cm⁻¹ region compared to pure HLW (Fig. 2a). We have to note that our measurements of reaction as well as DRIFTS were performed under conditions where water was not deliberately added to the gas streams. A recent report on an STM experiment demonstrated the importance of the film of water in CO₂ adsorption mechanism, eventually leading to formation of highly stable bicarbonates on rutile (110) (31). Furthermore, solvation effect of water decreases the energy barriers for CO₂ reduction and changes the selectivity of reaction processes on rutile (110) according to a recent first-principles calculation study (32).

A schematic representation of dimeric bicarbonate formation is given in Fig.4. Sorption takes place with initial interaction of CO_2 groups with hydroxyl groups on the surface. O-H groups make a nucleophilic attack to CO_2 , forming adsorbed bicarbonate structure. This kind of bicarbonate formation is consistent with the experimental observations indicating that the OH populations decrease upon CO_2 adsorption as well.

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	Table 2: Peak assignments	for CO ₂ adsorption	on TiO ₂ and	atase.
	Wavelength (cm ⁻¹)	Comments	Ref	This study
	3735,3725,3715	VOH	10,11,1 2	3690-3730
OH Stretching / Bending assignments	3690, 3675,3670,3665, 3640,3630(sh.)ª	vOH	10,11, 13,12	3660-3680
	3600-3200(br.)ª 3500-2800(br.)ª	_{voн} (surface and residual water)	13 13	3500-3000(br.)ª
	3350-3100(br.) ^a		10	
	1630, 1605	δон	10,13	1640
	3609, 3716		14	3728,3705, 3627,3600
CO ₂ Assignments	2375, 2360,2350, 2280 ^b		12	2360,2347, 2340,2335
	3340-3148, vон	M-bicarbonates ^c	15	
	2620-2450, vOH-O	D-bicarbonates ^c	15	2500-2700(br.)ª
	1702-1675,asym _{vC=0} 1672,1670	M-bicarbonates ^c Bidentate	15 13,16	
	1670(sh.) ^a	Bicarbonate	14	1682
	1655-1615,asym _{vC=0}	D-bicarbonates ^c	15	
	1630	Bicarbonate	13	1640
	1632-1600	CO ₃ ⁻ derivatives	10	
	1595,1578,1590-1575	Monodentate	13,16	
	1580	CO2 ⁻ derivatives	10	1550
Carbonate /	1555	Bicarbonate	14	
Bicarbonate Assignments	1410-1300	CO3 ⁻ derivatives	10	
Assignments	1400-1370,symm _{vC=0}	D-bicarbonates ^c	15	
	1430,1420,1408	Bicarbonate	14,13	1440
	1370-1320,1359,1315	Monodentate	13,16	1380
	1346-1327,symm _{vC=0} ,	M-bicarbonates ^c	15	
	1340	Bicarbonate	14	
	1320	CO2 ⁻ derivatives	10	1310
	1300,δ _{ОН-О}	D-bicarbonates ^c	15	
	(1252-1205,б _{он})	M-bicarbonates ^c	15	
	1243	Bidentate	13,16	
	1221,1220	Bicarbonate	14,13	1240
	1053	Bidentate	13	

^abr.:broad peak,sh.:shoulder,^{b 13}CO₂,^cOn metal oxides, M-monomeric, D-dimeric

Table 3: Carbonate bicarbonate region integrated DRIFTS intensities under vacuum, normalized with respect to their corresponding values under 500 Torr CO₂.^a Two pure HLW experiments are averaged.

Sample	Integrated Area (Vacuum / 500 torr CO ₂)
0.042 g/cm ² coat on grout	0.48
0.014 g/cm ² coat on grout	0.42
10% HLW in bulk grout	0.47
Pure HLW ^a	0.21
Pure grout	0.04

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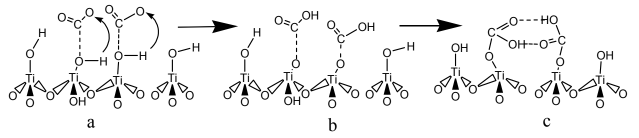


Fig. 4: Schematics of a) approach of CO₂ molecule to the surface, b) bicarbonate formation, c) dimeric bicarbonate formation.

CONCLUSIONS

The activity of the TiO_2 photocatalyst for benzene photooxidation declined by a factor of 3 when coated on a cementitious grout medium relative to the activity of a coating on a glass substrate. A detailed DRIFTS study unraveled the formation of

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stable carbonate-bicarbonates on TiO_2 surface in alkaline grout media. Also, dimeric bicarbonate interactions were observed on HLW-coated grout surfaces. We surmise that alkalinity in the grout can inhibit TiO_2 -active sites and decrease the photocatalytic activity.

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