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Organized Surface Functional Groups: Cooperative Catalysis via Thiol/Sulfonic Acid Pairing

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Abstract: The synthesis and characterization of heterogeneous catalysts containing surfaces functionalized with discrete pairs of sulfonic acid and thiol groups are reported. A catalyst having acid and thiol groups separated by three carbon atoms is ca. 3 times more active than a material containing randomly distributed acid and thiol groups in the condensation of acetone and phenol to bisphenol A and 14 times more active in the condensation of cyclohexanone and phenol to bisphenol Z. Increasing the acid/thiol distance in the paired materials decreases both the activity and selectivity. This work clearly reveals the importance of nanoscale organization of two disparate functional groups on the surface of heterogeneous catalysts.

Introduction

There is growing interest in developing multifunctional catalyst systems where each different functional group plays a distinctive role in the overall catalysis. Immobilization of two or more functional groups on a solid support allows for the possibility of spatial control of the different groups. They can be isolated from each other, which can be useful for sequential one-pot reactions, 1^{-4} or they can be intimately mixed such that direct interaction between the two groups is possible. In the latter case, the two groups can act in concert to provide activity greater than either could achieve alone (so-called cooperative catalysis).5-10

When seeking cooperative behavior between the functional groups, control of the distance between the reactive groups is essential in order to optimize the catalysis for a particular reaction. This is dramatically illustrated with enzymes that have multiple catalytic functional groups organized within a single active site. There have been several reports of the nanoscale arrangement of two or more *identical* functional groups on the

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surface of a silica catalyst,¹¹⁻¹⁵ but the arrangement of two different functional groups is more difficult. To our knowledge, there has been only one report of the organization of two disparate functional groups into pairs on a rigid support. Bass and Katz¹⁶ synthesized a precursor containing carbamate and xanthate groups that was grafted onto silica and subsequently thermolyzed to generate pairs of amine and thiol groups. No catalytic results were reported from this bifunctional solid.

One reaction where cooperative, heterogeneous catalysis has been well established is the synthesis of bisphenol A (BPA) from acetone and phenol. This reaction is catalyzed by strong acids and promoted by thiols that have been shown to increase both the yield and selectivity of p, p'-bisphenol A over the undesired o,p'-isomer (Scheme 1). Industrial processes typically employ sulfonated polystyrene resins promoted by thiols, such as cysteamine, bound to the resin by acid/base pairing.^{17,18} These resins have a random arrangement of acid and thiol groups, leading to a broad distribution of acid/thiol distances.

The mesoporous silica SBA-15¹⁹ is well suited as a support for immobilizing organized functional groups. Its large, uniform pore diameter (~ 6 nm) provides ample room for reactant and product diffusion, and its thick walls provide hydrothermal stability. The rigidity of the silica matrix ensures that the bound functional groups do not change their positioning (unlike polymeric supports that have the potential to shrink and swell

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Scheme 1. Condensation of Phenol with Acetone, Cyclohexanone, and Hexafluoroacetone To Form Bisphenols

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in different solvent environments). Furthermore, SBA-15 can be easily functionalized with a wide variety of functional groups by grafting an appropriate organosilane onto surface silanol groups in order to form covalent bonds to the surface.

We have recently shown²⁰ that SBA-15 functionalized with randomly distributed arylsulfonic acid and alkyl thiol groups is an effective catalyst for the synthesis of BPA. We also demonstrated that the acid and thiol groups must be near each other to achieve high activity and selectivity: a physical mixture of acid-functionalized and thiol-functionalized SBA-15 materials exhibited only moderate activity, whereas when the two functionalities were randomly distributed on the surface of the same SBA-15 material at high loadings (at a total organic loading $\sim 0.7 - 1.0$ mmol/g) the activity was nearly 4-fold higher, and the isomer ratio was 6-fold higher. We also showed that in this reaction the selectivity is essentially independent of conversion, which allows for the comparison of selectivities at different levels of conversion. We hypothesized that the thiol activates the protonated acetone molecule through formation of a propylidene sulfonium species that exhibits increased electrophilicity in addition to sterically disfavoring the o,p'isomer. If the acid and thiol groups are located near each other, the thiol can rapidly attack the protonated acetone molecule, generating the sulfonium species faster than if the groups are spatially isolated. Therefore we hypothesized that the catalytic activity should increase as the acid/thiol distance decreases.

Here, we report the synthesis of SBA-15 with alkylsulfonic acid and thiol groups organized into discrete pairs on the silica surface with varying distances between the acid and thiol groups. The distances are fixed by changing the nature of spacing groups between the acid and thiol functionalities, and the different spacers also provide the ability to tune both the acid/thiol distance and the electronic properties of the thiol. The resulting bifunctional catalysts give high activity and selectivity in the synthesis of bisphenol A. Pairing of the acid and thiol groups increases catalytic activity substantially compared to a randomly bifunctionalized catalyst.

Results and Discussion

Alkylsulfonic acid-functionalized mesoporous silica has been prepared through an oxidative method where surface-bound

propylthiol groups are oxidized with H2O214,21-25 or HNO326 to give propylsulfonic acid groups. Such an oxidative method is unsuitable for the preparation of bifunctional catalysts when the second functionality is also oxidizable. Furthermore, the oxidation is often incomplete and leads to disulfides and other partially oxidized species.^{22,25,26} To circumvent this difficulty we utilized a different route to alkylsulfonic acid groups based on a sultone intermediate. This intermediate serves both as a sulfonic acid precursor and as a site for anchoring a second functionality. Reaction of the sultone ring with a thiol-containing nucleophile generates the acid/thiol pair.

Sultone rings have been opened by a wide variety of nucleophiles, including halides,^{27–29} phosphines,^{30,31} thiolates,³² sulfides,³² alkoxides,^{27,33} and amines.^{33–35} In particular, if the nucleophile contains a thiol group, then the resulting material will contain pairs of acid and thiol groups.

In order to functionalize a silica surface with sultone moieties, the organosilane 1 was synthesized from 1,3-propanesultone and 3-iodopropyltriethoxysilane using a modification of the sultone alkylation method of Durst and Du Manoir.³⁶ Grafting of this silane onto SBA-15 in refluxing toluene resulted in the intermediate material SBA-g1 (Scheme 2). SBA-g1 is a versatile intermediate for generating bifunctional catalysts containing a

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Figure 1. ¹³C NMR (DMSO-d₆) of silane 1 (top) and ¹³C CP/MAS NMR of SBA-g1 (bottom). * denotes solvent peak.

Table 1	Catalyst	Characterization	Data
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entry	material	d (nm) ^a	S _{BET} (m ² /g) ^b	$D_p (nm)^c$	H ⁺ (mmol/g) ^d	S (mmol/g) ^e	S/H ⁺ (expected)	S/H+ (found)
1	SBA-15	11.3	810	6.1	N/A	N/A	N/A	N/A
2	SBA-g1	11.5	613	6.1	N/A	0.26	N/A	N/A
3	SBA-A	11.6	603	6.5	0.17	0.26	1	1.5
4	SBA-AS-p				0.38	0.81	2	2.1
5	SBA-AT-p	11.5	676	6.2	0.18	0.40	2	2.2
6	SBA-AT-r				0.18	1.04	N/A	5.8
7	SBA-AT-s1				0.22	0.54	3	2.5
8	SBA-AT-s2				0.18			
9	SBA-AT-s3				0.21			
10	SBA-AT-s4				0.17			
11	SBA-AT-s5	11.1			0.20			
12	SBA-AT-s6 ^f				0.16			
13	SBA-AT-s7		585	6.3	0.20	0.50	3	2.5
14	SBA-AT-s8				0.21	0.56	3	2.7

^{*a*} Unit cell parameter, from X-ray diffraction data. ^{*b*} Surface area, calculated from adsorption branch of N_2 isotherm using the BET method ^{*c*} Pore diameter from BJH analysis of desorption branch of N_2 isotherm. ^{*d*} From titration. ^{*e*} From elemental analysis (performed by QTI, Whitehouse NJ). ^{*f*} This material turned pink upon acidification due to a trace amount of organic byproduct resulting from the decomposition of the monosodium salt of *o*-xylyl dithiol.

sulfonic acid group and another functional group without the need for thiol oxidation.

SBA-g1 was characterized by cross-polarization magic angle spinning (CP/MAS) NMR, X-ray diffraction (XRD), nitrogen adsorption/desorption porosimetry, and elemental analysis. The ¹³C CP/MAS NMR spectrum shows the same resonances as the molecular precursor **1** (Figure 1); thus, the sultone ring remains intact throughout the grafting process. The sulfur content of the materials (from elemental analysis) was used as a measure of the organic loading, and it was found that approximately 30–80% of the silane used in the grafting reaction ended up covalently bound to the surface. The XRD pattern of the functionalized material is identical to that of the parent SBA-15 and indicates that long-range ordering of the material is not affected by the grafting process (see Supporting Information, Figure S2). Nitrogen adsorption/desorption porosimetry (Table 1, Figure S3) shows that the SBA-15 surface area drops from 810 to 613 m²/g upon grafting 1 at 0.2 mmol/g but that the pore diameter remains constant at 6.1 nm.

Sultone Ring-Opening. The sultone ring in **SBA-g1** can be hydrolyzed or thiolyzed to produce different surface groups (Scheme 3). Hydrolysis of the sultone in neutral water produces a solid that was used as a control catalyst because it contains acid groups but no thiols (**SBA-A**). Ring-opening with sodium hydrosulfide produces acid/thiol pairs separated by three carbon atoms (**SBA-AT-p**). ¹³C CP/MAS NMR (Figure 2) shows complete ring-opening of the sultone in both cases as evidenced by the disappearance of the carbon resonance at 68 ppm. The acid loadings of these materials were measured by ion exchange with NaCl followed by filtration and titration of the resulting HCl with NaOH.²⁰ The sulfur content of **SBA-AT-p** was found to be twice as high as the acid content measured by titration, confirming the 1:1 ratio of acid to thiol groups. The catalyst characterization data are summarized in Table 1.

Scheme 3. Ring-Opening of Surface-Bound Sultone To Form Sulfonic Acid-Containing Materials



Catalytic Reactions. The catalysts were used in the synthesis of bisphenol A. Surface loadings were purposely maintained low (\sim 0.2 mmol/g) in order to keep the resulting catalytic sites sufficiently isolated to observe the effects of discrete acid/thiol pairing. The results are given in Table 2. SBA-A gives a very low per-site-yield (PSY = 3.1, defined as mmol total products/ mmol H⁺) and selectivity (1.8, defined as p, p'/o, p' product ratio) due to the lack of thiol. Addition of homogeneous thiol improves both the yield and selectivity, although both are still only modest (10 and 8.4 respectively). SBA-AT-p is a highly active catalyst and gives a per-site-yield of 83 and a selectivity of 14. Addition of homogeneous thiol to this material decreases the per-siteyield to 74, and the selectivity is slightly increased to 15. A third catalyst was prepared via sultone thiolysis with sodium methanesulfide. This material contains pairs of acid and methyl sulfide groups, and is denoted SBA-AS-p. In the bisphenol A reaction SBA-AS-p is only slightly better than SBA-A (Table 2, Entry 5). These results confirm that the thiol functionality is necessary for high catalytic activity and also indicates that sulfide linkages can be used to tether other groups without substantially affecting catalysis (vide infra).

Homogeneous analogues of SBA-A and SBA-AT-p were also tested (Table 2, entries 6-8). 1-Propanesulfonic acid and 3-hydroxy-1-propanesulfonic acid gave nearly identical catalytic results. This suggests that the adjacent hydroxyl in SBA-A is not responsible for the poor catalytic activity of this material. The selectivity of these catalysts was similar to that of SBA-A, at a somewhat higher yield. 3-Mercapto-1-propanesulfonic acid also showed higher activity compared to that of its heterogeneous counterpart, SBA-AT-p, but somewhat lower selectivity. The reduced activity of the heterogeneous catalysts may be due to mass-transfer limitations, as the reagents must diffuse into the 1-D cylindrical pores of the SBA-15.

To investigate the possibility that the catalytic activity of SBA-AT-p was due to surface species leaching into solution, a hot-filtration test was performed. The solid catalyst was removed after 8 h, and no additional conversion was detected after an additional 9 or 16 h at 90 °C (see Figure 3). This result, along with the observation that unfunctionalized SBA-15 exhibits no catalytic activity toward bisphenol A, suggests that the catalytic activity of SBA-AT-p is entirely due to its surface-bound acid/ thiol pairs, rather than to the silica support or leached organic species.

Randomly-Distributed Acid/Thiol Catalysts. To test the effect of site organization, a catalyst containing randomly distributed alkylsulfonic acid and propylthiol groups was synthesized via the simultaneous grafting of two organosilanes. Silane 1 was again used as a source of alkylsulfonic acid groups since it allows for a nonoxidative synthesis. Since sultone rings can be opened by thiols at elevated temperatures, silane 2 was used as a source of disulfide-protected thiols. 1 and 2 were grafted onto SBA-15, followed by sultone hydrolysis and



Figure 2. Comparison of ¹³C CP/MAS NMR spectra of SBA-g1 (top) and sulfonic acid-containing SBA-15 materials. The complete disappearance of the sultone peak at 68 ppm indicates complete ring-opening of the sultone.

Table 2. Catalysis Data in Bisphenol A Synthesis^a for Heterogeneous and Homogeneous Catalysts

entry	heterogeneous catalyst	homogeneous catalyst	PSY ^b	isomer ratio ^c
1	SBA-A	none	3.1	1.8
2	SBA-A	PrSH	10	8.4
3	SBA-AT-p	none	83	14
4	SBA-AT-p	PrSH	74	15
5	SBA-AS-p	none	8.1	2.7
6	none	PrSO ₃ H	8.1	1.5
7	none	HOPrSO ₃ H	10	1.7
8	none	HSPrSO ₃ H	113	11

^{*a*} Reaction conditions: 0.02 mmol H⁺, 0 or 0.02 mmol propanethiol, 6 mmol acetone, 24 mmol phenol, 90 °C, 24 h. ^{*b*} Per site yield (mmol total product/mmol H⁺). ^{*c*} p,p'/o,p'.

disulfide reduction (Scheme 4). The resulting randomly distributed acid/thiol material is denoted **SBA-AT-r**. Due to the differential grafting efficiency of the two silanes, the acid/thiol ratio could not be controlled precisely at 1:1; elemental analysis showed that there were in fact nearly 5 times more thiol groups than acid groups. **SBA-AT-r** exhibited good selectivity in the synthesis of BPA, but with a nearly 3-fold reduction in persite-yield compared to that of the paired material **SBA-AT-p** (Table 3, entries 2–3).

To investigate the substrate scope of the heterogeneous catalysts, the ketone was varied to produce other bisphenol products. Using the more sterically hindered cyclohexanone as reactant (to produce bisphenol Z), the effect of pairing is even more pronounced (Table 3, entries 4-6). Thiol-free **SBA-A** gives low activity, the randomly bifunctionalized material gives only a small improvement, and the acid/thiol-paired catalyst has a much higher activity (14 times higher than for the randomly distributed catalyst). When hexafluoroacetone is used (to produce bisphenol AF), the effect of thiol promotion is greatly



Figure 3. Hot-filtration test results. Reaction conditions: 0.02 mmol H⁺, 6 mmol acetone, 24 mmol phenol, 90 °C. Catalyst was removed after 8 h, and filtrate was heated at 90 °C for another 16 h.

reduced, probably due to the greater reactivity of the fluorinated reactant (Table 3, entries 7–8). The increase in activity from **SBA-A** to **SBA-AT-p** is less than 2-fold. Thus, we conclude that the improvement in catalytic activity gained by pairing acid and thiol groups varies for different condensation reactions and is more pronounced for less reactive ketones.

Varying Acid/Thiol Distance. The acid/thiol distance was varied to investigate what effects separation distance has on activity and selectivity in catalyzing the synthesis of BPA. By opening the sultone ring with the monosodium salt of a dithiol, a thiol group is tethered to the acid via a sulfide linkage. In this way it is possible to tune both the length of the acid/thiol spacer and the electronic properties of the thiol (i.e., alkyl vs phenyl vs benzyl; see Scheme 3). ¹³C CP/MAS NMR showed quantitative conversion of the sultone moieties to sulfonic acids

Scheme 4. Simultaneous Grafting of Silanes 1 and 2 to Form Randomly Bifunctionalized Sultone/Disulfide Material and Subsequent Deprotection to Form Random Acid/Thiol Material SBA-AT-r



SBA-AT-r

Table 3. Catalysis Data for Heterogeneous Catalysts in Synthesis^a of Various Bisphenols

entry	heterogeneous catalyst	bisphenol product	PSY ^b	isomer ratio ^c
1	SBA-A	А	3.1	1.8
2	SBA-AT-p	А	83	14
3	SBA-AT-r	А	29	22
4	SBA-A	Z	0.3	N/A^d
5	SBA-AT-p	Z	14	13
6	SBA-AT-r	Z	1.0	N/A^d
7	SBA-A	AF	8.4	N/A^d
8	SBA-AT-p	AF	15	N/A^d

^{*a*} Reaction conditions: 0.02 mmol H⁺, 6 mmol ketone, 24 mmol phenol, 90 °C, 24 h. Catalyst loading ~0.2 mmol H⁺/g. ^{*b*} Per site yield (mmol total product/mmol H⁺). ^{*c*} *p*,*p*'/*o*,*p*'. ^{*d*} *o*,*p*'-isomer below detection limit.

for these products (Figure S6, S7). Elemental analysis revealed sulfur loadings to be approximately three times the acid loading (Table 1, Entries 7, 13, 14), corresponding to the expected 1:1 acid/thiol ratio, and the surface area, pore size, and long-range ordering were maintained. The resulting materials with various alkyl and aryl spacers (denoted SBA-AT-s1 through SBA-ATs8) were used to catalyze the BPA reaction (Table 4). Catalysts SBA-AT-s1, -s2, and -s3 (ring-opened by 1,2-ethanedithiol, 1,3propanedithiol, and 1,6-hexanedithiol, respectively) showed reduced activity and selectivity compared to SBA-AT-p, where the acid and thiol groups are in closer proximity (Table 4, entries 1-4). The activity decrease with distance is quite dramatic, as increasing the length of the acid/thiol spacer by three atoms (SBA-AT-s1 vs SBA-AT-p) decreases the yield by a factor of 2. Additional increase in the spacer length by one atom (SBA-AT-s2) reduces yield by another factor of 2, down to a level similar to that of SBA-AT-r (Table 3, entry 3). SBA-AT-s3 exhibits activity similar to that of SBA-AT-s2.

Catalysts **SBA-AT-s4** and **-s5**, containing *ortho-* and *para* phenylthiol groups, respectively, are both poor catalysts (Table 4, entries 5–6). In particular, the *ortho* variant shows activity

 Table 4.
 Catalysis Data in Bisphenol A Synthesis^a for

 Heterogeneous Acid/Thiol Spacer Catalysts

	heterogeneous			
entry	catalyst	spacer type	PSY [♭]	isomer ratio ^c
1	SBA-AT-p	none	83	14
2	SBA-AT-s1	alkyl, $n = 2$	42	12
3	SBA-AT-s2	alkyl, $n = 3$	20	6.5
4	SBA-AT-s3	alkyl, $n = 6$	22	6.8
5	SBA-AT-s4	aryl, $m = 0$, ortho	5.2	1.9
6	SBA-AT-s5	aryl, $m = 0$, para	21	4.3
7	SBA-AT-s6	aryl, $m = 1$, ortho	64	9.6
8	SBA-AT-s7	aryl, $m = 1$, meta	37	7.9
9	SBA-AT-s8	aryl, $m = 1$, para	47	7.9

^{*a*} Reaction conditions: 0.02 mmol H⁺, 6 mmol acetone, 24 mmol phenol, 90 °C, 24 h. Catalyst loading ~0.2 mmol H⁺/g. ^{*b*} Per site yield (mmol total product/mmol H⁺). ^{*b*} *p*,*p*'/*o*,*p*'.

no better than that of the thiol-free material **SBA-A**. The poor activity of this material could be due to steric hindrance between the adjacent sulfide and thiol groups that could prevent the thiol from reacting with acetone. The reduced nucleophilicity of phenyl thiols vs alkyl thiols is also likely responsible for some of the lower activity of these two catalysts.

Catalysts **SBA-AT-s6**, **-s7**, and **-s8** (derived from *o*-, *m*-, and *p*-xylyl dithiols, respectively) contain benzyl thiols. These materials generally show greater activity than those containing alkyl and phenyl spacers (Table 4, entries 7–9). The catalyst **SBA-AT-s6**, containing an *o*-benzyl spacer is the most active of all the spacer materials investigated here. We believe this is because the thiol is positioned such that it can point directly at the acid group and shortens the acid/thiol distance.

Conclusions

The distance between the acid and thiol group in a series of heterogeneous catalysts was shown to have a profound influence on the catalytic behavior in the synthesis of bisphenols. These data highlight the importance of the spatial positioning of functional groups in the development of heterogeneous catalysts and also reveal how the nanoscale organization of catalytic surfaces can be tuned to provide a level of reactivity control unachievable through traditional random functionalization approaches.

In the case of acid and thiol groups, it was found that the catalytic activity for bisphenol synthesis is highest when the two groups are as close as possible. It may be that for other pairs of groups, such as mutually destructive ones, that the activity vs distance behavior could be qualitatively different.

The sultone-containing silica **SBA-g1** was used here to generate acid/thiol pairs. By the reaction of this intermediate with other nucleophiles, it is possible to create a wide array of surface-paired materials.

Experimental Section

Materials. Tetrahydrofuran (THF) and toluene were distilled over sodium immediately before use. All other solvents were analytical grade and used as received. 3-Iodopropyltriethoxysilane was prepared from 3-chloropropyltriethoxysilane (Gelest) and sodium iodide according to the literature procedure.³⁷ Organosilane **2** was synthesized from 3-mercaptopropyltriethoxysilane and 2,2'-dipyridyldisulfide according to the literature procedure¹⁴ followed by purification by chromatography

⁽³⁷⁾ Matsura, V.; Guari, Y.; Larionova, J.; Guerin, C.; Caneschi, A.; Sangregorio, C.; Lancelle-Beltran, E.; Mehdi, A.; Corriu, R. J. P. J. Mater. Chem. 2004, 14, 3026–3033.

on silica gel. Anhydrous sodium hydrosulfide was purchased from Alfa Aesar. All other chemicals were purchased from Aldrich and used as received. All reactions were performed under an argon atmosphere.

3-(3-Triethoxysilylpropyl)-[1,2]-oxathiolane 2,2-dioxide (Organosilane 1). 1,3-Propanesultone (2.7 g, 22 mmol) was dissolved in dry THF (50 mL). The solution was cooled to -78 °C, and *n*-butyllithium (1.6 M in hexanes, 14 mL, 22 mmol) was added slowly over 10 min. After stirring for another 10 min, 3-iodopropyltriethoxysilane (3.3 g, 10 mmol) was added slowly over 5 min. After stirring for 15 min, the temperature was raised to -60 °C. After stirring for 9 h, the temperature was raised to 0 °C, water (50 mL) was added, and the mixture was transferred to a separatory funnel. The aqueous layer was removed, and the organic layer was washed with saturated NaCl solution (50 mL) and dried over anhydrous MgSO4. After the solvent was removed in vacuo, chromatography on silica gel (3:2 hexanes/ethyl acetate, $R_{\rm f}$ = 0.3) afforded 1 (0.61 g, 19%) as a colorless liquid. ¹H NMR (300 MHz, DMSO- d_6) δ 4.38 (m, 2H), 3.73 (q, J = 7 Hz, 6H), 3.48 (m, 1H), 2.58 (m, 1H), 2.13 (m, 1H), 1.70 (m, 2H), 1.46 (m, 2H), 1.13 (t, J = 7 Hz, 9H), 0.61 (t, J = 8 Hz, 2H). ¹³C NMR (300 MHz, DMSO d_6) δ 68.6, 58.4, 55.5, 31.6, 29.9, 20.6, 18.9, 10.3. Anal. Calcd for C12H26O6SSi: C, 44.15; H, 8.03; S, 9.82. Found: C, 43.05; H, 7.46; S, 9.85. HRMS (FAB⁺): *m*/*z* (M + H⁺) (C₁₂H₂₇SiSO₆) Calcd, 327.1298; Found, 327.1302.

SBA-g1 (Sultone-Functionalized Silica). SBA-15 (1.0 g, synthesized according to the literature procedure³⁸) was dried under flowing argon at 170 °C for 4 h. After cooling, dry toluene (50 mL) was added via syringe, and the mixture was stirred vigorously to form a uniform suspension. An appropriate amount of **1** (typically ~0.7 mmol) was added dropwise via syringe. The suspension was stirred for 45 min at room temperature and then refluxed for 16 h. After cooling to room temperature, the solids were filtered and washed with toluene and dichloromethane (3 × 20 mL each). The solids were Soxhlet extracted with dichloromethane overnight, dried under vacuum, and stored under an argon atmosphere in a drybox until further use.

SBA-A (Thiol-Free Sulfonic Acid Catalyst). SBA-g1 (0.5 g) was added to water (40 mL), and the resulting suspension was stirred at 40 °C for 2 days. Then the solids were filtered, washed with water (3 \times 30 mL), 0.5 N HCl (3 \times 30 mL), and water (4 \times 30 mL), and dried under high vacuum overnight.

SBA-AT-p (Acid/Thiol-Paired Catalyst). **SBA-g1** (0.5 g) was suspended in anhydrous DMF (15 mL). Anhydrous sodium hydrosulfide (86 mg, 1.5 mmol) was dissolved in anhydrous DMF (10 mL), and the resulting blue solution was added dropwise to the stirred silica suspension. After stirring for 24 h at room temperature, the solids were filtered, washed with DMF (3×30 mL) and water (5×30 mL), and then suspended in 0.5 N HCl (30 mL) to acidify. After stirring for 3 h the white solids were filtered and washed with 0.5 N HCl (3×30

(38) Zhao, D. Y.; Huo, Q.; Feng, J.; Chmelka, B.; Stucky, G. J. Am. Chem. Soc. 1998, 120, 6024–6036. mL) and water (5 \times 30 mL) and dried under high vacuum overnight. **SBA-AS-p** was prepared with the same procedure, except that sodium methanesulfide was used in place of sodium hydrosulfide.

SBA-AT-r (Random Acid/Thiol Catalyst). SBA-15 (0.5 g) was grafted with 1 (0.16 g, 0.5 mmol) and 2 (0.17 g, 0.5 mmol) according to the procedure above for **SBA-g1**. The two organosilanes were added dropwise simultaneously. After Soxhlet extraction and drying, the palebeige solids were suspended in water (50 mL) and stirred at 40 °C for 2 days. The solids were filtered, washed with water (2×30 mL) and 0.5 N HCl (3×30 mL), and suspended in an aqueous solution of tris(2-carboxyethyl)phosphine hydrochloride (0.015 M, 50 mL). This suspension was stirred at 55 °C for 2 days, and then the solids were filtered, washed with water and methanol (4×30 mL each), and dried under high vacuum overnight to yield a nearly white powder.

SBA-AT-s (Acid/Thiol Spacer Catalysts). Dithiol (15 mmol) was dissolved in anhydrous DMF (10 mL), and sodium hydride (60 wt % in mineral oil, 5 mmol) was added. This mixture was stirred until all solids had dissolved and then was added slowly to a suspension of **SBA-g1** (0.5 g) in anhydrous DMF (10 mL). After stirring for 24 h at room temperature, the solids were filtered, washed, and acidified as described for **SBA-AT-p**.

Acid Titration. To the silica material to be analyzed (\sim 30 mg) was added aqueous NaCl (2 N, 4 mL), and the suspension was stirred for 24 h. The solids were removed by filtration and washed with water (4 \times 2 mL), and the combined filtrate was titrated with 0.01 N NaOH using phenol red as indicator.

Catalytic Reaction: Condensation of Phenol and Ketone. An amount of catalyst corresponding to 0.02 mmol H⁺ (~100 mg) was added to a vial and dried under high vacuum at 80 °C overnight. Phenol (2.2 g, 24 mmol) and ketone (6 mmol) were added, and the vial was sealed under argon and stirred at 90 °C for 24 h. The catalyst was removed by filtration and washed with acetonitrile to a total filtrate volume of 25 mL, and the products were quantified by HPLC. Persite-yield was calculated on the basis of the number of acid sites present, and selectivity was defined as the ratio of bisphenol isomers (p,p'/o,p').

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Supporting Information Available: X-ray diffraction data, ¹H NMR spectrum of silane 1, nitrogen adsorption/desorption isotherms, ²⁹Si CP/MAS spectrum of **SBA-g1**, ¹³C CP/MAS NMR spectra of **SBA-AT-r** and spacer materials **SBA-AT-s**(1– 8). This material is available free of charge via the Internet at http://pubs.acs.org.

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