



Palladium-Catalyzed Aerobic Dehydrogenation of Substituted Cyclohexanones to Phenols Yusuke Izawa *et al. Science* **333**, 209 (2011); DOI: 10.1126/science.1204183

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tional period, which is on the order of picoseconds. $E_{\rm s}$ between high- and low-chalcocite phases results from the different Cu arrangements in these two phases and the related extra Ewald energy at the interface. This is similar to the case of the interface between wurtzite (WZ) and zincblende (ZB), where the different second nearest-neighbor atomic positions cause different Ewald energies in these two phases. It has been found that the interface energy between WZ and ZB per surface unit cell is similar to the energy difference per unit cell (28). Thus, if we take this approximation that the interface energy between low- and high-chalcocite is the same as the internal energy difference between these two phases, $(\varepsilon_1 - \varepsilon_2) = 40$ meV per unit cell (23), and assume that there are $N \text{ Cu}_2\text{S}$ unit formulae inside a spherical core, we have $E_s =$ $(36\pi)^{1/3}N^{2/3}(\varepsilon_1 - \varepsilon_2)$. Note that $C = C_{\text{unit}}N$, and $C_{\text{unit}} = 52 \text{ J/mol} \cdot \text{K}$ (23). Thus, from Eq. 3, we get $\tau \sim 2$ s when N = 1000 and $\tau_0 = 1$ ps (29). This fluctuation time is of the same order as our observed experimental value.

In summary, we have observed dynamic structural transformations of a single Cu₂S nanorod from a low- to a high-chalcocite structure. The influence of the surface and interface energies on nucleation and pinning phenomena of a particular phase by defects suggests strategies for stabilizing metastable structures. The ability to directly visualize these processes will aid in the future design of materials with new and controlled phases.

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Supporting Online Material

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Palladium-Catalyzed Aerobic Dehydrogenation of Substituted Cyclohexanones to Phenols

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Aromatic molecules are key constituents of many pharmaceuticals, electronic materials, and commodity plastics. The utility of these molecules directly reflects the identity and pattern of substituents on the aromatic ring. Here, we report a palladium(II) catalyst system, incorporating an unconventional *ortho*-dimethylaminopyridine ligand, for the conversion of substituted cyclohexanones to the corresponding phenols. The reaction proceeds via successive dehydrogenation of two saturated carbon-carbon bonds of the six-membered ring and uses molecular oxygen as the hydrogen acceptor. This reactivity demonstrates a versatile and efficient strategy for the synthesis of substituted aromatic molecules with fundamentally different selectivity constraints from the numerous known synthetic methods that rely on substitution of a preexisting aromatic ring.

Phenols are common precursors and core structures of industrial chemicals ranging from pharmaceuticals to polymers. The introduction of chemical functional groups with specific patterns around the aromatic ring represents a key challenge in the preparation of these molecules (1). Electrophilic aromatic substitutions are classical chemical reactions that remain among the most versatile methods for the synthesis of substituted phenols; however, strong electronic directing effects associated with these reactions limit their utility to the preparation of ortho- and para-substituted derivatives. This limitation has inspired extensive efforts to identify complementary routes to substituted phenols, such as a recent

two-step arene C-H borylation/oxidation procedure for the introduction of a hydroxyl group into an aromatic ring, guided by steric rather than electronic effects (2). Recent advances in palladium-catalyzed aerobic oxidation reactions (3-5) suggested to us that diverse phenol derivatives, including those with meta substitution, could be accessed by dehydrogenation of cyclohexanones via sequential Pd-mediated C-H activation/\u03c3-hydride elimination steps, followed by tautomerization of the resulting dienone product (Fig. 1A). This strategy is appealing because PdII-hydride intermediates formed in this mechanism could be oxidized by molecular oxygen (6, 7), thereby enabling the overall process to be catalytic in Pd with water as the sole by-product (Fig. 1B). Successful catalysts for this class of reactions could find broad utility owing to the numerous straightforward chemical reactions that provide access to substituted cyclohexanones, including enolate arylation and alkylation methods, conjugate addition to cyclohexenones, and Robinson annulation and Diels-Alder reactions (Fig. 1C).

The preparation of phenols from ketone precursors have been explored previously (8–16). Condensation reactions of acyclic ketones, for example, with β -ketoaldehydes or β -diketones, enable direct access to substituted phenols (8), but low product yields, limited access to starting materials, and/or formation of isomeric products

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have restricted the utility of these procedures. Methods for formation of phenols via dehydrogenation of cyclohexenones have been pursued (8-13), but reactions of this type typically use undesirable stoichiometric reagents, such as DDQ (2,3-dichloro-5,6-dicyano-1,4-benzoquinone) (17); use stepwise procedures, such as bromination/ dehydrobromination; or require harsh reaction conditions (≥200°C) that limit functional group compatibility. In contrast, no effective methods for dehydrogenation of substituted cyclohexanones exist, with relevant precedents almost exclusively limited to reactions of unsubstituted cyclohexanone and yields of $\leq 30\%$ (mostly $\leq 5\%$) (12–16). Iridium complexes bearing tridentate "pincer" ligands are among the most effective catalysts for the dehydrogenation of saturated hydrocarbons (18, 19). These reactions are typically carried out in the presence of a sacrificial hydrogen acceptor, such as norbornene or tert-butylethylene, and a recent investigation of the reaction of an Ir-pincer complex with cyclohexanone resulted in stoichiometric dehydrogenation. Catalytic turnover was inhibited by the formation of an Irphenoxide product, (pincer)Ir(H)(OPh) (20).

In order to explore prospects for the proposed Pd-catalyzed dehydrogenation methods, we investigated the reactivity of 3-methylcyclohexenone (1a) and 3-methylcyclohexanone (1b) under 1 atm of O2 with various PdII catalysts. Preliminary analysis of Pd^{II} sources, solvents, and reaction conditions revealed that 3-methylphenol (meta-cresol) could be obtained in modest yields from 1a and 1b (28 to 52%) with 3 mol % Pd(OAc)₂ or Pd(TFA)₂ (OAc is an acetate group and TFA is trifluoroacetate) in dimethylsulfoxide (DMSO) as the solvent under 1 atm of O₂ at 80°C (Fig. 2, entries 1 and 2) (21, 22). Brønsted bases, such as sodium acetate, and traditional monodentate and bidentate nitrogen ligands (pyridine, bipyridine, and phenanthroline) failed to increase the yield of the phenol product (entries 3 to 6; for expanded screening results, see table S2). The use of 4,5diazafluorenone (23) led to a substantial increase in the yield of *m*-cresol from 1a (78%, entry 7); however, conversion to phenol from the saturated cyclohexanone 1b was still unsatisfactory. Both key steps in the substrate oxidation sequence, C-H activation and β-hydride elimination (Fig. 1A), should benefit from a more electrophilic catalyst, and recent success with 2-fluoropyridine (^{2F}py) as an electron-deficient ligand in aerobic oxidative coupling reactions (24) prompted us to evaluate ligands of this type in the dehydrogenation reactions (entries 8 to 15). The use of ^{2F}py as a ligand led to a modest improvement in the yield of 2 from 1b (44%, entry 10); however, the best results were obtained when 2-(N,N-dimethylamino) pyridine (^{2NMe2}py) was used in combination with p-toluenesulfonic acid (TsOH) (entry 18). We speculate that the improved results obtained with the ^{2NMe2}py/TsOH combination relative to those with ^{2NMe2}py alone (compare entries 15 and 18) reflect the ability of TsOH to protonate the tertiary amine of the coordinated pyridine

and thereby afford a more electron-deficient pyridine ligand. The position of the dimethylamino group on the pyridine ligand is important, as revealed by the inferior results obtained with 4-(N,N-dimethylamino)pyridine in place of the 2-substituted ligand (entries 19 and 20). Common heterogeneous palladium catalysts failed to afford the desired *m*-cresol (entries 21 to 24).

The optimized catalytic conditions (Fig. 2, entry 18) proved to be effective in the preparation of a number of substituted phenol derivatives (Fig. 3). Varying the position of the methyl substituent on the cyclohexanone had little effect on the outcome of the reaction; the corresponding ortho-, meta-, and para-cresols were each obtained in good yield (Fig. 3 entries 1 to 3). Aryl-substituted phenols, including a number of meta-substituted derivatives, were accessed via the dehydrogenation of the corresponding arylcyclohexanone derivatives (entries 4 to 11). The 3-arylcyclohexanones were readily obtained via conjugate addition of aryl boronic acids to cyclohexenone, and both electron-donating and electron-withdrawing groups were tolerated in the dehydrogenation reaction. Substrates with aryl groups bearing a para bro-

mide or iodide afforded only low yields of the desired phenol (28% and 16%, respectively; not shown in Fig. 3). Diels-Alder cycloaddition and Robinson annulation represent classical and widely used synthetic organic reactions that provide efficient routes to cyclohexenones bearing multiple substituents on the six-membered ring. The aerobic oxidation method described here provides an attractive alternative to the use of stoichiometric reagents, such as DDQ, which have been used previously in the dehydrogenation of cyclohexenones (17), and such substrates underwent very effective dehydrogenation under the optimized catalytic conditions, including those bearing alkyl, aryl, and/or ester substituents (entries 12 to 17).

The 3,5-disubstituted cyclohexenones were pursued further, because the corresponding phenol derivatives exhibit important biological activity and products of this type cannot be readily prepared by classical aromatic-substitution, metal-catalyzed cross-coupling, or related synthetic methods. As a representative example, *O*-terphenylcarbamate **5** was recently identified as a potent in vitro allosteric inhibitor of the human luteinizing hormone



Fig. 1. Strategy for the synthesis of phenols via aerobic oxidative dehydrogenation of cyclohexanone derivatives. (**A**) Stepwise sequence for Pd-mediated dehydrogenation of cyclohexanone. (**B**) Catalytic mechanism whereby cyclohexanone dehydrogenation can be achieved with O_2 as the terminal oxidant. (**C**) Representative synthetic methods that afford facile access to substituted cyclohexanone derivatives. L, a ligand; M, metal; R, organic substituent; X, halide or pseudohalide.

receptor, which has been implicated in fertility and ovarian cancer (25). This compound was prepared by traditional Suzuki coupling methods

Fig. 2. Catalyst optimization for aerobic oxidative dehydrogenation of cyclohexanone derivatives 1aand 1b. Reaction conditions are as follows: cyclic ketone (1.0 mmol), PdX₂ (0.03 mmol), ligand/TsOH (mol % indicated), DMSO (0.4 ml), 80°C, 1 atm O₂, 24 hours. Et, ethyl group; Me, methyl group; TsOH, *p*-toluenesulfonic acid. Yields determined by gas chromatography. using 3,5-dibromophenol as the starting material; however, introduction of the unsymmetrical aryl substitution pattern results in low yields of the biaryl intermediate 6 and the desired 3,5diarylphenol 4 (30% and 62% yields, respectively; Fig. 4). In contrast, the 3,5-diarylcyclohexenone



[†]Yield in parentheses based on 5 mol % Pd.



* 3% Pd/6% ligand/12% TsOH. [†] 5% Pd/10% ligand/20% TsOH.

Fig. 3. Palladium-catalyzed aerobic dehydrogenation of cyclic ketones. Reaction conditions are as follows: cyclic ketone (1.0 mmol), Pd(TFA)₂ (0.03 or 0.05 mmol), 2-(*W*,*N*-dimethylamino)pyridine (0.06 or 0.10 mmol), TsOH (0.12 or 0.20 mmol), DMSO (0.4 ml), 1 atm O₂, 80°C, 24 hours. Isolated product yields are reported.



Fig. 4. Application of palladium-catalyzed aerobic oxidative dehydrogenation in the preparation of a terphenyl-derived allosteric inhibitor of the luteinizing hormone receptor. Ph, phenyl group.

derivative **3** is obtained readily from very inexpensive starting materials (4-methylacetophenone, benzaldehyde, and acetone) via sequential aldol condensation and Robinson annulation. Subsequent Pd-catalyzed dehydrogenation of **3** afforded phenol **4** in excellent yield.

Preliminary mechanistic analysis of these dehydrogenation reactions was performed by monitoring the conversion of cyclohexanone to phenol by gas chromatography. The kinetic time course revealed the formation and disappearance of the partially dehydrogenated intermediate, cyclohexenone (Fig. 5A). This result is consistent with the overall catalytic mechanism in Fig. 1B, in which the substrate dissociates from the catalyst after each dehydrogenation step. A fit of the kinetic data based on a simple sequential reaction model reveals that the two dehydrogenation steps have similar rate constants, $k_1 = 0.12(\pm 0.02)$ hour⁻¹ and $k_2 = 0.33(\pm 0.04)$ hour⁻¹ (Fig. 5B). The dehydrogenation of cyclohexenone, monitored independently, exhibits a rate constant somewhat lower than that obtained from the fit of the sequential reaction, $k_2' = 0.19(\pm 0.02)$ hour⁻¹. Accurate interpretation of these results will require further investigation; however, the higher concentration of cyclohexenone in the latter reaction may slow the catalytic turnover via alkene coordination to Pd.

The joint application of catalyst screening and mechanistic studies will play an important role in extending these reactions to different product classes. For example, catalysts that can effect the first step substantially more rapidly than the second step $(k_1:k_2 > 10:1)$ would enable selective formation of the enone products rather than phenols. Moreover, it should be possible to develop efficient catalysts for dehydrogenative aromatization of other substrate classes, such as substituted cyclohexenes (26-28). Substrates of this type are readily accessed via Diels-Alder cycloaddition reactions, and their dehydrogenation could proceed via sequential allylic C-H activation/β-hydride elimination steps (Fig. 5C). Toward this end, preliminary studies reveal that cyclohexene derivative 7 undergoes efficient dehydrogenation to the trisubstituted arene 8 in near-quantitative yield.



Fig. 5. (**A**) Kinetic profile of Pd^{II}-catalyzed aerobic dehydrogenation of cyclohexanone and cyclohexenone, showing the formation and decay of cyclohexenone as an intermediate in the reaction. Reaction conditions are as follows: cyclohexanone (0.5 mmol), Pd(TFA)₂ (0.025 mmol), 2-(*N*,*N*-dimethylamino)pyridine (0.05 mmol), TsOH (0.10 mmol), DMSO (0.5 ml), 1 atm O₂, 80°C. Error bars represent standard deviations from five independent measurements. (**B**) Comparison of the rate constants obtained for the dehydrogenation steps in the sequential conversion of cyclohexanone to phenol and in the direct dehydrogenation of cyclohexenone. (**C**) General strategy and a specific example of Pd-catalyzed aerobic dehydrogenation of cyclohexenes.

Methods for selective dehydrogenation of saturated carbon-carbon bonds represent an important class of C–H functionalization (18, 29), and the reactions presented above highlight the prospective utility of such methods in the synthesis of substituted aromatic molecules. These reactions achieve high conversions and product yields, they are capable of using O_2 as the hydrogen acceptor,

and the catalyst tolerates useful substrate functional groups, including aromatic and heteroatom substituents. With the development of improved methods for safe and scalable aerobic oxidation reactions (30), dehydrogenation methods of this type could have an important impact on laboratoryand industrial-scale chemical synthesis.

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Supporting Online Material

www.sciencemag.org/cgi/content/full/science.1204183/DC1 Materials and Methods Figs. S1 to S20 Tables S1 and S2 Characterization Data of New Compounds References (31–43)

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High Pre-Eruptive Water Contents Preserved in Lunar Melt Inclusions

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The Moon has long been thought to be highly depleted in volatiles such as water, and indeed published direct measurements of water in lunar volcanic glasses have never exceeded 50 parts per million (ppm). Here, we report in situ measurements of water in lunar melt inclusions; these samples of primitive lunar magma, by virtue of being trapped within olivine crystals before volcanic eruption, did not experience posteruptive degassing. The lunar melt inclusions contain 615 to 1410 ppm water and high correlated amounts of fluorine (50 to 78 ppm), sulfur (612 to 877 ppm), and chlorine (1.5 to 3.0 ppm). These volatile contents are very similar to primitive terrestrial mid-ocean ridge basalts and indicate that some parts of the lunar interior contain as much water as Earth's upper mantle.

The Moon is thought to have formed in a giant impact collision between a Marssized object and an early-formed proto-Earth (1). Though all of the inner planets, including Earth, are depleted in water and other volatiles when compared with primitive meteorites, the more extreme depletion of volatiles in lunar volcanic rocks has long been taken as key evidence for a giant impact that resulted in high-temperature catastrophic degassing of the material that formed the Moon (2, 3). However, recent work on rapidly quenched lunar volcanic glasses has detected the presence of water dissolved in lunar magmas at concentrations up to 46 parts per million (ppm) (4), and water contents of lunar apatite grains from mare basalts are consistent with similarly minor amounts of water in primitive lunar magmas (5-7). These results indicate that the Moon is not a perfectly anhydrous planetary body and suggest that some fraction of the Moon's observed depletion in highly volatile elements may be the result of magmatic degassing during the eruption of lunar magmas into the nearvacuum of the Moon's surface.

To bypass the process of volcanic degassing, we conducted a search for lunar melt inclusions in Apollo 17 sample 74220, a lunar soil containing ~99% high-Ti volcanic glass beads, the so-called orange glass with 9 to 12 weight % (wt %) TiO₂ (8). Melt inclusions are small samples of magma trapped within crystals that grow in the magma before eruption. By virtue of their enclosure within their host crystals, melt inclusions are protected from loss of volatiles by degassing

during magma eruption. Melt inclusions have been used for decades to determine pre-eruptive volatile contents of terrestrial magmas from subduction zones (9, 10), hotspots (11, 12), and midocean ridges (13, 14), as well as volatile contents of martian magmas (15, 16). Using standard petrographic methods, we identified nine inclusionbearing olivine crystals (Fig. 1) and analyzed melt inclusions hosted within (17). The measured water contents of the melt inclusions range from 615 ppm to a maximum of 1410 ppm (Fig. 2); these water contents are up to 100 times as high as the water content of the matrix glass surrounding the olivine crystals (6 to 30 ppm H₂O) and the centers of individual volcanic glass beads from the same sample (4). The melt inclusions also contain high concentrations of fluorine (50 to 78 ppm), sulfur (612 to 877 ppm), and chlorine (1.5 to 3.0 ppm) that are 2 to 100 times as high as those of the matrix glasses and individual glass beads from this sample (Fig. 3). Volatile contents corrected for postentrapment crystallization are on average 21% lower than the measured concentrations (17) and represent the best estimate of the pre-eruptive concentrations of volatiles in the 74220 magma.

There are few descriptions in the literature of melt inclusions contained within olivine from lunar samples, but these existing observations provide important context for the volatile abundances we have observed. Roedder and Weiblen (18-20) noted the presence of silicate melt inclusions in the first samples returned from the Apollo 11, Apollo 12, and Luna 16 missions and reported that many primary melt inclusions contained a vapor bubble, requiring dissolved volatiles to have been present in the melt at the time it was trapped within the host crystal. Klein and Rutherford (21) and Weitz *et al.* (22) found sulfur contents of 600 to 800 ppm, similar to our

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