

Titanium and Zinc Oxide Nanoparticles Are Proton-Coupled Electron Transfer Agents Joel N. Schrauben *et al. Science* **336**, 1298 (2012); DOI: 10.1126/science.1220234

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Supplementary Materials

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Titanium and Zinc Oxide Nanoparticles Are Proton-Coupled Electron Transfer Agents

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Oxidation/reduction reactions at metal oxide surfaces are important to emerging solar energy conversion processes, photocatalysis, and geochemical transformations. Here we show that the usual description of these reactions as electron transfers is incomplete. Reduced TiO_2 and ZnO nanoparticles in solution can transfer an electron and a proton to phenoxyl and nitroxyl radicals, indicating that e^- and H⁺ are coupled in this interfacial reaction. These proton-coupled electron transfer (PCET) reactions are rapid and quantitative. The identification of metal oxide surfaces as PCET reagents has implications for the understanding and development of chemical energy technologies, which will rely on e^-/H^+ coupling.

xidation/reduction reactions of metal oxide materials (MO_x) play a key role in emerging energy technologies, as well as other processes such as photocatalytic wastewater remediation and self-cleaning surfaces (I)and geochemical redox formation and dissolution of minerals (2). These interfacial (solid/solution) redox processes are generally described just in terms of electron transfer (ET). For instance, the extensive literature on photochemical and electrochemical water splitting by MO_x is focused on the ET between surface-bound species and the valence and conduction bands of the solid. In dye-sensitized solar cells (DSSCs), the excited state of a dye injects an e^- into nanocrystalline TiO_2 (3). All of these disparate processes are affected by proton activity (the pH of aqueous interfaces), indicating a role for H⁺. Here, we report well-characterized examples of reduced TiO₂ and ZnO nanoparticles (TiO₂/e⁻ and ZnO/e⁻) reacting by proton-coupled electron transfer [PCET (4)]. TiO₂ and ZnO have been extensively studied from both fundamental and technological

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Fig. 1. Pictures of asprepared and reduced (A) ZnO and (B) amorphous TiO₂ nanoparticles in toluene. UV-visible spectra of solutions of (C) ZnO (20µM particles, 3.9-nm diameter) and (D) amorphous TiO_2 (51 μ M, 3 nm) during irradiation with a 200-W Hg/Xe lamp. ^tBu₃ArO[•] titrations of (E) ZnO/e-(0.51 mM, 3.9 nm) after 30 min of irradiation and (F) TiO_2/e^- (51 µM, 3 nm) after 10 min of irradiation.



perspectives [nearly 58,000 publications on TiO₂ alone (5)], and well-characterized nanoparticle systems have been described. The demonstration that these reduced oxides can transfer both electrons and protons expands the traditional view of interfacial redox reactions. The occurrence of interfacial PCET brings together previous electrochemical, photochemical, and photocatalytic reports of H⁺/e⁻ coupling in TiO₂ (1, 3, 6–9) and ZnO (10), as discussed below. PCET is likely a common theme for MO_x reactivity in the presence of protons (11, 12).

Dodecylamine-capped ZnO nanocrystals (3 to 6 nm in diameter) and oleic acid–capped amorphous and anatase TiO₂ particles (3 and 5 nm in diameter, respectively) were prepared and characterized by well-known procedures (13-15). Ultraviolet (UV) irradiation of these materials forms electron/hole pairs (excitons), and rapid quenching of the holes by oxidation of organic species at the surface (1) leaves extra electrons in the particles. Toluene solutions of these ZnO/ e^- and

TiO₂/ e^- particles, as previously reported, are longlived in the absence of air; are blue or purple/ black, respectively, because of optical and nearinfrared absorbances (Fig. 1, A to D); and have characteristic electron paramagnetic resonance (EPR) spectra (9, 16–19). The added electrons occupy delocalized, conduction band-like orbitals in ZnO/ e^- (19, 20), whereas in TiO₂/ e^- they are much more localized on a few Ti atoms (11).

Addition of the stable 2,4,6-tri-*tert*-butylphenoxyl radical (^{*I*}Bu₃ArO[•]) to air-free toluene solutions of ZnO/ e^- or TiO₂/ e^- (amorphous or anatase) yields the oxidized particles and the corresponding phenol ^{*I*}Bu₃ArOH, as indicated by ¹H nuclear magnetic resonance (NMR), EPR, optical spectroscopies, and gas chromatography (GC) (Fig. 2A). A similar reaction is observed for irradiated anaerobic slurries of Aeroxide P25 TiO₂ (Evonik Industries AG, Essen, Germany), a commercially available anatase with average particle diameter ~20 nm, so this result is not limited to organic-capped soluble nanoparticles. ZnO/ e^- and TiO₂/ e^-



Fig. 2. (**A**) The ¹H NMR spectra in toluene- d_8 of (top) ^{*t*}Bu₃ArOH, (middle) amorphous TiO₂ (50 µM, ~3 nm, irradiated for 30 min) and ^{*t*}Bu₃ArO[•] (45 equivalents per particle), and (bottom) the corresponding reaction for ZnO (2.5 equivalents ^{*t*}Bu₃ArO[•], 1.9 mM particles, ~4 nm, irradiated for 30 min), showing formation of ^{*t*}Bu₃ArOH (circles) (asterisks indicate solvent peaks; peaks of the particle capping groups are unlabeled). (**B**) Optical spectra of as-prepared TiO₂ particles (black), TiO₂/e⁻ (red), and ^{*t*}Bu₃ArO[•] (blue); the wavelengths of the exciting laser (pump) and monitoring position (probe) are indicated. (**C**) Typical kinetic traces monitoring the absorbance at 620 nm as a function of time after the laser flash: black points, TiO₂ only; red points, TiO₂ and ^{*t*}Bu₃ArO[•] (0.78 mM) with fit to second-order kinetics (supplementary material pp. 7 and 8, figs. S10 to S14, and table S1). (**D**) Plot of second-order rate constants *k* [e.g., from (**C**)] versus concentration of ^{*t*}Bu₃ArO[•].

are also oxidized by the nitroxyl radical 2,2,6,6tetramethyl-piperidin-1-yl-oxyl (TEMPO) to give the reduced hydroxyl amine TEMPOH. The organic radicals do not react with nonreduced particles, particles that have not been irradiated, or particles that have been reduced and then reoxidized by air. These reactions (Eqs. 1 and 2) occur in toluene, a solvent without acidic protons. Therefore, the formation of 'Bu₃ArOH from ^tBu₃ArO[•], and TEMPOH from TEMPO, must involve transfer of e^- and H^+ from the particles to the organic radical; the proton is explicitly shown in the balanced Eqs. 1 and 2. Reduced TiO₂ has previously been shown to accomplish multielectron/ multiproton reductions, but often the solvent (typically water) was assumed to provide the protons (9, 17, 18).

$$H^{+}-TiO_{2}/e^{-} + XO^{\bullet} \rightarrow TiO_{2} + XOH$$
(1)
$$H^{+}-ZnO_{2}/e^{-} + XO^{\bullet} \rightarrow ZnO_{2} + XOH$$
XO[•] = 'Bu₂ArO[•], TEMPO (2)

The active protons likely come both from the particle syntheses and from the photochemical charging. The particles were prepared hydrolytically and were not calcined, so surface hydroxyl groups are undoubtedly present. The particles are capped by organic ligands (vide supra) and likely contain residual solvents from syntheses, particularly ethanol. Analyses of ZnO/e⁻ solutions by GC and GC-mass spectroscopy (GC-MS) show that quenching of the photolytically formed hole yields one primary product: the imine from condensation of dodecylamine and acetaldehyde, the latter from $2e^{-}/2H^{+}$ oxidation of ethanol (Eq. 3, R = dodecyl). The identity of the product was confirmed by independent synthesis. Thus, protons are formed commensurate with the electrons in the reduced particles. (No photocharging products derived from the toluene solvent have been observed.)

$$2h^+$$
 + CH₃CH₂OH + H₂NR \rightarrow CH₃CH=NR +
H₂O + 2H⁺ (3)

Titrations of ZnO/e^- solutions show a gradual disappearance of the ZnO/e⁻ absorbance and then, after the endpoint, appearance of the spectrum of the excess 'Bu₃ArO[•] (Fig. 1E). Addition of a small excess of ^tBu₃ArO[•] causes disappearance of the ZnO/e^{-} resonance in the EPR spectrum and appearance of the signal for the remaining ^tBu₃ArO[•]. Titrations with the pure electron acceptor decamethylferrocenium (Cp*₂Fe⁺) are very similar, and titrations of different aliquots of the same ZnO/e^{-} solution with $Cp*_{2}Fe^{+}$ and ^tBu₃ArO[•] give the same "concentration of electrons" within the $\pm 10\%$ uncertainty of the titrations (from repetitive measurements; fig. S4). Thus, all of the spectroscopically visible conduction band (CB) electrons in ZnO/e⁻ can be removed either as e^- , with Cp*₂Fe⁺, or as e^- accompanied by H^+ with ^tBu₃ArO[•]. The titration results, together with the total metal concentration and the average particle diameter, give the average number of electrons per particle (n_{e^-}) ; 3.9-nm ZnO

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particles irradiated for 30 min typically show $n_{e^-} \approx 4$, consistent with prior work (19).

Titrations of TiO2/e- solutions behave similarly, giving the same value of n_{e^-} within error with use of either 'Bu3ArO' or ferrocenium (Cp_2Fe^+) (fig. S5). The derived extinction coefficient per electron ε_{e^-} (700 nm) = 633 ± 70 M^{-1} cm⁻¹ is similar to that in other reduced TiO₂ systems (18) ($\pm 1\sigma$ error, from the data in fig. S5). Irradiation of the amorphous particles was found to cause extensive reduction, $n_{e^-} \approx 45$ per 3-nm particle after 1 hour based on the above ε_{e^-} . Continued irradiation results in slower additional reduction, up to $n_{e^-} \approx 72$ after 10 hours. These values correspond, respectively, to reduction of 10% and 16% of all the Ti^{4+} ions to Ti^{3+} . Anatase TiO2 nanocrystals are photochemically reduced to a smaller extent, ~2% Ti³⁺ ($n_{e^-} \approx 40$ per 5-nm nanocrystal) after >4 hours of irradiation. For comparison, the nanostructured TiO₂ in an operating dye-sensitized solar cell is ~0.03% reduced (usually reported as $\sim 10^{18} e^{-1}$ /cm³) (21), and the ZnO/e^- particles above could be considered 0.3% reduced. The greater reduction of the TiO_2 particles is likely due in part to the localized nature of the reduced sites, and the very extensive reduction of the amorphous TiO₂ is attributed to the ability of the charge-balancing protons to be close to the added electrons.

The kinetics of the PCET reactions have been interrogated with flash photolysis using ~5-ns pulses of 355-nm light. With amorphous TiO₂ particles in the absence of an oxidant, irradiation causes an increased absorbance at 620 nm within ~10 ns that does not decay over several ms (Fig. 2, B and C, black points), indicating the formation of a stable TiO_2/e^- solution (Fig. 1C) (18, 22, 23). When the particles are excited in the presence of ${}^{t}Bu_{3}ArO^{\bullet}$, this feature is observed transiently but decays on a ~200-us time scale to an absorbance lower than the initial value (Fig. 2C, red points). The decay is due to consumption of both TiO_2/e^- and ${}^tBu_3ArO^{\bullet}$. Kinetic data over a range of concentrations of ^tBu₃ArO[•] and TiO₂ particles follow secondorder kinetics and yield the bimolecular rate constant $(1.0 \pm 0.4) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ (this value is the average $\pm 1\sigma$ for many measurements between 0.1 and 2.0 mM ^tBu₃ArO[•] with data from two different batches of particles; Fig. 2D and table S1). Reactions of the anatase particles occur with a similar rate constant of $(1.5 \pm 0.4) \times$ $10^7 \text{ M}^{-1} \text{ s}^{-1}$ (from six measurements between 0.2 and 0.7 mM 'Bu₃ArO[•]; fig. S10). These are fast reactions, within $\sim 10^3$ of the estimated diffusion limit (see supplementary materials, p. 9). Oxidations of amorphous TiO_2/e^- with the pure electron acceptors methylviologen or Cp*₂Fe⁺ proceed even faster (> 10^{7} M⁻¹ s⁻¹, fig. S11). The kinetics of ZnO/e⁻ and ^tBu₃ArO[•] were measured similarly, monitoring the recovery of the bleached ZnO band edge. Reactions of two different batches of ZnO/e⁻ particles with ^tBu₃ArO[•] gave rate constants of (3.0 \pm 0.6) \times 10 7 M^{-1} s^{-1} and (6.9 \pm 1.3) \times $10^{6} \text{ M}^{-1} \text{ s}^{-1}$ (from 13 measurements between 0.5

and 2 mM 'Bu₃ArO[•], fig. S12). When an excess of the dodecylamine capping group was added to the latter batch, *k* was reduced by nearly a factor of two (table S1). This suggests that the oxidant must reach the surface of the particle and that different dodecylamine surface coverage may explain the variation in *k* between batches of particles. The reactions of TEMPO are much slower than those of 'Bu₃ArO[•]: Amorphous TiO₂/*e*⁻ + TEMPO has a second-order rate constant just onethousandth as high $[(1.1 \pm 0.5) \times 10^4 \text{ M}^{-1} \text{ s}^{-1}]$ (from six measurements at 25 mM TEMPO, fig. S13), and for one batch of ZnO particles (fig. S14) the rate constant for reaction with TEMPO was nearly a factor of 40,000 smaller.

These PCET reactions are unlikely to proceed by a mechanism of initial outer-sphere ET, even though this is the common proposal for interfacial redox reactions. The reduction of TEMPO to its anion is in particular thermodynamically very difficult (24), and the formation of either ^tBu₃ArO or TEMPO- is unfavorable in the nonpolar toluene solvent used. ^tBu₃ArO[•] or TEMPO could bind to a coordinatively unsaturated surface metal center, followed by protonation to give free XOH. This would be analogous to nitroxyl radicals binding to molecular Ti³⁺ complexes such as Cp₂Ti^{III}Cl to form Ti⁴⁺-nitroxide species (25–27) (Cp = η^5 -C₅H₅). However, a high density of unsaturated titanium centers seems unlikely given the hydrolytic synthesis of the particles and the tightly held oleate capping ligands. In addition, XO[•] binding would be expected to be more rapid for the less sterically encumbered TEMPO radical, but this radical reacts 10³ to 10⁴ times slower than ^tBu₃ArO[•]. The reactions thus more likely proceed by a concerted proton-electron transfer (CPET), equivalently called hydrogen atom (H[•]) transfer (HAT). Both ^tBu₃ArO[•] and TEMPO are common H[•] abstractors (24). The faster reactions of 'Bu₃ArO[•] are consistent with a HAT mechanism because the O-H bond formed by ^tBu₃ArO[•] is 11.5 kcal mol⁻¹ stronger than that formed by TEMPO (24). With use of our model of bimolecular HAT rate constants (k_{HAT}) based on the Marcus cross relation (28), this greater driving force should be associated with a $\sim 10^4$ faster k_{HAT} , in good agreement with the measured values. These reactions of metal oxide particles are therefore analogous to HAT from molecular Ti³⁺-aquo complexes to carbon radicals R[•] (Eq. 4) (29).

$$Cp_2(Cl)Ti^{III}(OH_2) + R^{\bullet} \rightarrow Cp_2(Cl)Ti^{IV}(OH) + R-H$$

$$k = 1.0 \times 10^5 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$$
 (4)

The demonstration here that MO_x particles can react by PCET provides an expanded view of interfacial charge transfer reactions and the involvement of protons. From an energetic perspective, proton activity (pH in aqueous solutions) is known to strongly modulate the effective CB energy (E_{CB}) of bulk metal oxides. Although this behavior has traditionally been ascribed to changes in surface protonation (30), recent work indicates that each electron added to the oxide is

accompanied by a proton (3, 6-8, 31-34). Most notably, Hupp and co-workers showed proton uptake upon electrochemical reduction of TiO2 (6, 7). This is analogous to the addition of both e^{-} and Li⁺ upon charging of a lithium battery anode (6, 33, 35) and to H'-doped ZnO and TiO_2 (1, 9, 10). These and other studies indicate that reduced and protonated metal oxides are formed under a variety of electrochemical and chemical conditions, not only by the photochemical charging route used here. The thermochemical coupling between E_{CB} and pH is related to the $e^{-/H^{+}}$ (PCET) reactivity described here, and together they have broad implications. Transfer of e^- and H^+ to or from a surface requires at least a partially localized description, in contrast to the pure delocalized band-structure pictures typically used for interfacial ET and for PCET reactions in which interfacial ET is coupled to proton movements within a solute or solvent (36-38). For instance, the PCET reducing power of the particles studied here reflects their ability to donate e^{-} and H⁺ together, not just their ability to donate electrons, and is thus only indirectly related to the E_{CB} or Fermi energy (just as the potential of a lithium battery anode is not the $E_{\rm CB}$ of the oxide). The ability to transfer e^- and H⁺ together should be advantageous in metal oxide catalysis, electrocatalysis, and photocatalysis of reactions such as water oxidation $(2H_2O \rightarrow$ $O_2 + 4H^+ + 4e^-$), because moving the two particles together can avoid high-energy chemical intermediates (4).

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Supplementary Materials

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Interglacial Hydroclimate in the Tropical West Pacific Through the Late Pleistocene

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Records of atmospheric carbon dioxide concentration (Pco_2) and Antarctic temperature have revealed an intriguing change in the magnitude of interglacial warmth and Pco_2 at around 430,000 years ago (430 ka), but the global climate repercussions of this change remain elusive. Here, we present a stalagmite-based reconstruction of tropical West Pacific hydroclimate from 570 to 210 ka. The results suggest similar regional precipitation amounts across the four interglacials contained in the record, implying that tropical hydroclimate was insensitive to interglacial differences in Pco_2 and high-latitude temperature. In contrast, during glacial terminations, drying in the tropical West Pacific accompanied cooling events in northern high latitudes. Therefore, the tropical convective heat engine can either stabilize or amplify global climate change, depending on the nature of the climate forcing.

The study of past interglacial climates provides valuable insight into natural climate variability under conditions roughly similar to those of the present day, but with differences in atmospheric carbon dioxide concentration $(Pco_2)(1)$ and astronomic forcing. Especially intriguing is a step-like increase in interglacial Antarctic temperature (2), global ice melting (3), and Pco_2 (4, 5) that took place at roughly 430,000 years ago (430 ka) and is referred to as the Mid-Brunhes Event (MBE). To understand the underlying cause of this shift toward warmer interglacials, it is crucial to establish the global signature of interglacial climates during that time, but sufficiently long records, especially from low latitudes and terrestrial archives, are sparse. Because Pco2 and ice cover affect the planet's radiative balance, a global change in interglacial climate at the MBE might be expected. Although many marine sea surface temperature (SST) reconstructions from around the globe show qualitatively similar changes to those observed in Antarctica (6), the few available lower-latitude SST reconstructions (7–9) from this time period do not show the same consistent increase in post-MBE interglacial temperatures. Given that many models simulate strong dynamical links between high-latitude and tropical climate [e.g., (10)], more tropical records are required to test and improve our understanding of low-latitude climate sensitivity to a range of different forcing factors.

Here, we present stalagmite records from northern Borneo (4°N, 115°E; fig. S1) that span several glacial-to-interglacial cycles and include the MBE. We reconstruct Warm Pool hydroclimate from the oxygen isotope (δ^{18} O) variability recorded in three stalagmites with overlapping growth periods. Borneo lies in the West Pacific Warm Pool, which is an important heat and moisture source for higher latitudes (*11*). Although there is little seasonality in modern rainfall amount at our site, the isotopic composition of precipitation varies seasonally by up to 2 to 3‰ (*12*), with the most enriched δ^{18} O values occurring in February and March, when the intertropical convergence zone (ITCZ) is furthest south of our site. However, on interannual time scales, the El Niño– Southern Oscillation (ENSO) phenomenon dominates both precipitation amount and δ^{18} O, with El Niño events resulting in large-scale drying and rainfall δ^{18} O that is roughly 5 to 6‰ higher than average (*12*). Two recent studies of water isotopes in the same area conclude that regional (much more than local) rainfall amount is the main driver for rainfall δ^{18} O variability on intraseasonal and longer time scales today (*13*) and during modeled millennial-scale events in the past (*14*).

We analyzed stalagmites from three different caves in Gunung Mulu and Gunung Buda National Parks, each about 5 to 10 km apart. The stalagmites were dated by $^{238}\mathrm{U}\text{-}^{234}\mathrm{U}\text{-}^{230}\mathrm{Th}$ measurements (15) and cover the time period from 570 (±20) ka to 210 (±10) ka [from marine isotope stage (MIS) 13 to the beginning of MIS7], with some gaps due to hiatuses. Sample GC08 covers the whole time period, whereas samples Squeeze1 and WR5 yield shorter records that overlap GC08. In comparison to other cave systems, dating of these stalagmites is challenging due to their unusually low ²³⁴U/²³⁸U ratios and/or low U concentrations (fig. S4 and tables S1 to S3). The old age of the samples exacerbates these dating issues. Furthermore, there is evidence that slight open-system exchange of U has affected sample GC08, requiring us to model the ages in this stalagmite by assuming a constant initial $^{234}\text{U}/^{238}\text{U}$ ratio (15). Varying detrital 230 Th contamination cannot explain the age reversals in this stalagmite, owing to the short half-life of 230 Th compared to the age of our samples (15). The error assigned to the assumed constant initial ²³⁴U/²³⁸U ratio leads to increased age uncertainty in the GC08 chronology, with individual age errors ranging from ± 9200 to $\pm 13,500$ years (2σ ; table S1), limiting our ability to compare the millennial-scale timing of climate change in Borneo with other records. The age constraints are sufficient, however, to identify each interglacial stage, which is the main focus of this study. In sample WR5, open-system alteration might also have occurred but cannot be corrected for (15). The δ^{18} O record from this stalagmite therefore

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