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Molecularly rigid porous polyamine host enhances barium titanate catalysed H₂O₂ generation†

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Barium titanate (BTO) is well-known (as a photo- or sono/piezo-catalyst) to produce hydrogen peroxide via 2-electron reduction of oxygen in the presence of a sacrificial guencher, such as isopropanol. While barium titanate nanoparticles with a tetragonal crystal structure (piezoelectric) are particularly reactive, the recovery and reuse of these nano-catalysts from reactions can be difficult. Here, barium titanate nanoparticles of typically 200 nm to 600 nm diameter are embedded into a host film of a polymer of intrinsic microporosity (PIM-EA-TB). Due to molecular rigidity of the polymer, there is no capping effect, and the surface catalytic reaction occurs effectively with a catalyst embedded in the polymer. In this exploratory work, the catalytic formation of H2O2 in the presence of isopropanol is investigated via kinetic studies and by electron paramagnetic resonance (EPR). Perhaps surprisingly, at a neutral pH the rate of the catalytic reaction is substantially increased when barium titanate is embedded into the polymer host and when the polymer is protonated. This is attributed here to a "kinetic cage effect" which exploits the tertiary amine in the polymer backbone with anodic and cathodic processes coupled into a pH neutral reaction.

1. Introduction

Barium titanate (BTO) is a phase-changing ferroelectric material.¹ Barium titanate has been employed as a catalyst in a range of potential applications; this has included piezo-photocatalysis,² water pollution treatment,3,4 hydrogen production and antibacterial coatings,⁵ in hydrogen peroxide formation and therapeutic applications,6 and for bio-piezoelectric platforms.7 In most cases, the catalytic process is suggested to be associated with piezoelectric effects,8 namely it is linked to mechanically induced/enhanced electric fields at crystal surfaces that lower the activation barrier. The activation of barium titanate has been achieved by light,9 by ultrasound,10 or even by mechanical stirring. 11 Barium titanate is synthesised in the form of powders, or nanopowders, and is often simply added to the reaction. Here, the use and recovery of the barium titanate catalyst is attempted by embedding the catalyst nanopowder into a molecularly rigid microporous host polymer (PIM-EA-TB). In this regard, nanocomposites based on barium titanate in polymer hosts are of considerable interest. 12 Perhaps surprisingly, an enhanced catalytic reactivity is observed.

Polymers of intrinsic microporosity^{13,14} (PIMs) have emerged over the past decade with intriguing properties due to a molecularly rigid structure enhancing processability (the poor intermolecular contact increases solubility) and inducing intrinsic microporosity with typically 1-2 nm diameter pores. 15 A prototypical PIM is PIM-EA-TB¹⁶ ("EA" for ethanoanthracene and "TB" for Tröger base, see molecular structure in Fig. 1) with a typical BET surface area of 939 m² g⁻¹ and a pore volume of approx. $0.9 \text{ m}^3 \text{ g}^{-1}$, depending on the degree of protonation.¹⁷ The tertiary amine has been observed to be protonated at approx. pH 4. The molecularly rigid backbone of the polymer avoids nanoparticle surface capping¹⁸ and ensures full accessibility and reactivity of the embedded nanocatalysts. 19 In fact, cages can form at the interface where the polymer can affect the catalytic reaction. Examples for applications of PIMs in catalysis include embedded photocatalysts for hydrogen generation,20

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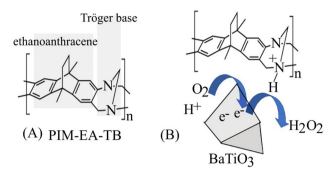


Fig. 1 (A) Molecular structure of PIM-EA-TB. (B) Illustration of BaTiO₃ catalyst reacting to produce hydrogen peroxide when embedded into PIM-FA-TR

catalysis for hydrogen peroxide production,21 and molecular catalysis for alcohol oxidation.²² Usually, the PIM acts as an inert host in a similar way to zeolites or other inert host materials. However, in a recent report an example was presented whereby the use of PIM-EA-TB enhanced the catalytic conversion of formate oxidase activity.²³ It was suggested that in a twostep process the reduction of oxygen to hydrogen peroxide was enhanced by protonation of the tertiary amine within the PIM-EA-TB cages. A similar catalytic system is investigated here based on barium titanate.

The reduction of barium titanate leads to blue Ti(III) radical colour centres as observed by EPR spectroscopy²⁴ and by optical spectroscopy. In addition, barium titanate exists in distinct phases²⁵ due to temperature dependent symmetry-breaking of the high-temperature cubic perovskite phase and lowtemperature tetragonal phase. The resulting interfacial strain and electric fields have been utilised in catalysis. In addition to pure barium titanate, cerium-doped barium titanate26 and silver nanoparticle modified barium titanate materials²⁷ have been employed in enhanced piezo- and photocatalysis. Here, we demonstrate that barium titanate reactivity is enhanced by embedding into a porous polymer matrix.

There have been previous reports of barium titanate immobilisation into siloxane foams^{28,29} and into PVDF³⁰ for water treatment applications. Here, barium titanate nanoparticles are investigated bare and when embedded into PIM-EA-TB. For the case of hydrogen peroxide production, it is demonstrated that the catalytic surface chemistry is retained and the polymerembedded nanopowder is easily recovered and reused. A catalysis enhancing effect of the PIM-EA-TB cavity on hydrogen peroxide production is observed in acidic or neutral solution and assigned to enhanced dioxygen reduction.

2. Experimental

2.1. Chemical reagents

PIM-EA-TB $(C_{21}H_{20}N_2)_n$ with 1.08 g cm⁻³ was synthesised following a literature procedure.31 Chloroform (CHCl3, 1.49 g cm^{-3} , $119.38 \text{ g mol}^{-1}$, Sigma-Aldrich, >99% purity), isopropanol (C₃H₈O, 0.78 g cm⁻³, 60.1 g mol⁻¹, Sigma-Aldrich, >99% purity), dimethyl sulfoxide ((CH₃)₂SO₄, 1.1 g cm⁻³,

78.13 g mol⁻¹, Sigma-Aldrich, > 99% purity), para-nitrophenyl boronic acid (C₆H₆BNO₄, 1.4 g cm⁻³, 166.93 g mol⁻¹, Sigma-Aldrich, >99% purity), sodium bicarbonate (NaHCO₃, 2.2 g cm^{-3} , $84.007 \text{ g mol}^{-1}$, Sigma-Aldrich, > 99% purity) were obtained commercially and used without further purification.

2.2. Barium titanate synthesis and characterisation

Barium titanate (BTO, BaTiO₃) was synthesised following a literature procedure³² with an annealing step at 1000 °C; this annealing temperature led to maximum H₂O₂ production when compared to lower or higher annealing temperatures. Scanning electron microscopy images were recorded using a Jeol JSM-7900F field emission scanning electron microscope. The SEM images Fig. 2(a) and (b) data (Fig. 2) show particles of 0.37 \pm 0.10 μm diameter. Transmission electron microscope images were recorded using a Jeol JEM-2100PLUS TEM. The fringe pattern in Fig. 2(d) is consistent with the (011) pattern dominating as observed previously and the transmission electron microscopy (TEM) fringe pattern is consistent with the (011) pattern dominating as observed previously.33 The FFT pattern suggests a separation of 3.54 nm⁻¹ which suggests a lattice constant of 3.99 Å. The XRD pattern in Fig. 2E is closely matching literature data^{34,35} for tetragonal BaTiO3 (P4mm). The surface area of the particles measured with BET surface area 5.82 m² g⁻¹ (Fig. 2F).

2.3. Immobilising barium titanate by PIM-EA-TB

An amount of 0.01 g barium titanate, a white crystal powder was combined 1:1 with a 2 mL solution of PIM-EA-TB (0.01 g) in chloroform. After 15 minutes of sonication, the suspension was drop-cast onto a 1×1 cm² filter paper (Whatman no. 1). The rapid evaporation of chloroform resulted in a uniform layer of PIM-EA-TB with embedded BaTiO₃, thereby forming an immobilized and easily recoverable/re-usable catalyst. Fig. 3 shows SEM images of the porous composite.

2.4. Quantification of H₂O₂ production

Hydrogen peroxide production and detection involved the reaction of 1 mg BaTiO3, that was either (i) bare or (ii) embedded in PIM-EA-TB@BaTiO3, with a 90% deionized water and typically 10% isopropanol solution. After a 5 minute reaction time, the sample solution was filtered or the catalyst removed, and the hydrogen peroxide concentration was quantified by mass spectroscopy indirectly via reaction with para-nitroboronic acid to paranitrophenol (see ESI†).36 Mass spectrometry (LC-MS) analyses were performed on an Automated Agilent QTOF (Walkup) used with HPLC (4 chromatography columns) and variable wavelength detector based on an Agilent QTOF 6545 with a Jetstream ESI† spray source coupled to an Agilent 1260 Infinity II Quat pump HPLC with a 1260 autosampler, and a column oven compartment and variable wavelength detector.

2.5. Electron paramagnetic resonance (EPR)

Electron paramagnetic resonance (EPR) spectroscopy measurements were performed on a Bruker Elexsys E500 spectrometer equipped with a Bruker ER4122-SHQE X band resonator. EPR spectra were recorded at either room temperature or 98 K and

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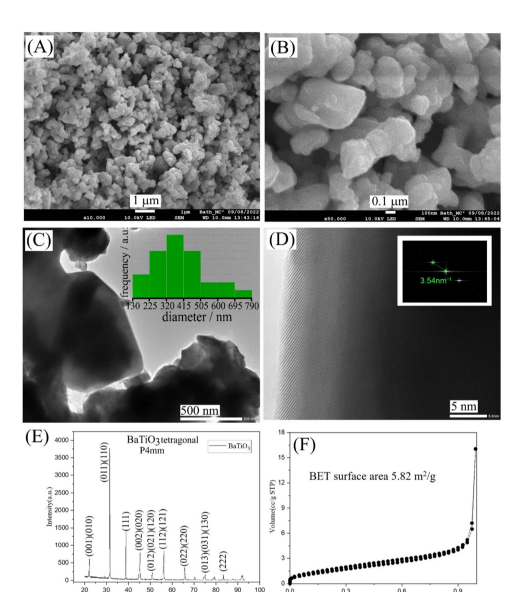


Fig. 2 (A) and (B) SEM images of BaTiO₃ particles with typically 0.37 ± 0.10 µm diameter. (C) and (D) TEM images showing BaTiO₃ aggregates and fringe pattern (Inset: Fourier transform with lattice constant 3.54 nm⁻¹.). (E) XRD pattern. (F) Nitrogen (BET) adsorption isotherm.

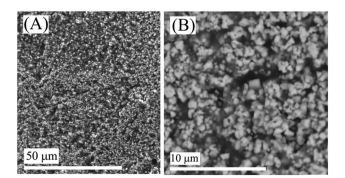


Fig. 3 (A) and (B) SEM images of BaTiO₃ particles (white) embedded into PIM-EA-TB (grey) in 1:1 weight ratio.

100 kHz magnetic field modulation frequency, 0.4 mT magnetic field modulation amplitude, 10.24 ms conversion time, 60 dB receiver gain, 4096 number of points, and 10 averages per spectrum. The powder samples were degassed overnight at 393 K under a dynamic vacuum in a Schlenk line prior to EPR measurements. EPR spectra were simulated using the Easyspin toolbox for Matlab.37

Results and discussion

3.1. Contrasting reactivity of bare and immobilised barium titanate

Initially, reactivity tests were performed with 1 mg BaTiO₃ catalyst suspended in 1 cm3 water (containing 10 vol% isopropanol) with gentle agitation. Fig. 4A (red symbols) shows the production of H₂O₂ with time (thermocatalytic). An increase in

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H₂O₂ concentration occurs over the first 30 minutes with a plateau probably due to simultaneous consumption of H₂O₂. Next, the same amount of catalyst powder immobilised within a PIM-EA-TB host polymer was deposited onto a cellulose substrate and placed into 1 cm³ water (containing 10 vol% isopropanol) with only natural convection. Fig. 4A (black symbols) shows that the production of H2O2 under these conditions essentially doubles, despite using the same conditions and the same amount of catalyst. The process continues up to about 60 minutes, then reaching a plateau close to 0.3 mM H₂O₂. The plateau is likely to be associated with the competing reduction of H₂O₂ to water. Next, by purging the solution with pure oxygen (replacing ambient oxygen) the same reaction (Fig. 4A, blue symbols) leads to the formation of approx. 5 times more H₂O₂. That is, the process appears to be first order in oxygen. The mechanism can be written in two sequential reaction steps (eqn (1) and (2).

Isopropanol
$$\rightleftharpoons$$
 acetone + 2H⁺(aq) + 2e⁻(barium titanate)
(1)

$$O_2(aq) + 2H^+(aq) + 2e^-(barium titanate) \rightleftharpoons H_2O_2(aq)$$
 (2)

The mechanism is assigned to a surface process based on (i) isopropanol being oxidised and the transfer of two electrons into the conduction band (essentially interfacial Ti(III)),

(ii) dioxygen from solution interacting with the BaTiO₃ surface and transfer of two electrons to form H₂O₂. This kind of process with a slow oxygen reduction reaction could be facilitated by the presence of protonated tertiary ammonium site on the PIM-EA-TB polymer backbone (see Fig. 1B) to explain the observed increase in catalytic rate. However, the pH within the polymeric structure is not well defined, and it might switch between two states during the course of the reaction cycle within the polymeric cages containing the barium titanate catalyst (vide infra). Next, the effect of the amount of catalyst was investigated. Fig. 4B shows good linearity with the amount of catalyst building up substantial concentrations of H₂O₂ for both the bare barium titanate catalyst and the PIM-EA-TB embedded catalyst. The rate of formation of H₂O₂ appears to be first order in BaTiO3 surface sites. With the BaTiO3 embedded into PIM-EA-TB, the rate of H₂O₂ production approximately doubles in all cases. The PIM-EA-TB polymer provides a microporous environment for effective diffusion of reagents. The molecular rigidity of the polymer prevents blocking of the catalyst surface sites.

The investigation of the effect of the amount of isopropanol is shown in Fig. 4C. When doubling the concentration of isopropanol from 10 vol% to 20 vol%, the rate of H₂O₂ production doubles. Therefore, the rate of H₂O₂ formation is first order in isopropanol for both the bare barium titanate catalyst

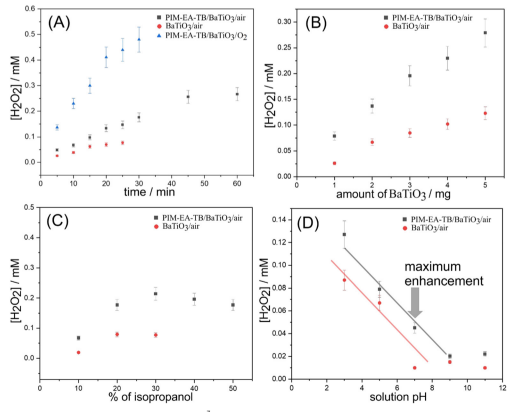


Fig. 4 (A) Production of H₂O₂ with time for 1 mg BaTiO₃ in 1 cm³ water with 10 vol% isopropanol (square: with PIM-EA-TB; circle: pure catalyst; triangle: in pure oxygen). (B) Effect of the amount of catalyst in water with 10 vol% isopropanol; 5 min. (C) Effect of vol% isopropanol on H_2O_2 production for 1 mg BaTiO₃: 5 min. (D) Production of H₂O₂ with 1 mg BaTiO₃ in 1 cm³ water with 10 vol% isopropanol as a function pH (adjusted by adding very small quantities of aqueous NaOH or HCl). Errors estimated at $\pm 10\%$.

and PIM-EA-TB embedded catalyst. Only with very high concentrations of isopropanol of 30 vol% or higher is a plateau, or a decline, in H₂O₂ production observed. This is consistent with a process that is first order in isopropanol and saturation of surface sites. The presence of isopropanol at the catalyst surface could also slow down the oxygen reduction to result in this behaviour.

The effect of the solution pH (for 10 vol% isopropanol) is shown in Fig. 4D. Perhaps interestingly, the reaction does not work at all in the alkaline pH range (with or without polymer host). Only at a pH 6, or more acidic, is BaTiO3 effective for oxygen reduction and H2O2 production. This observation matches with the reported point zero charge (pzc) at approx. pH 4.38 Protonation of the surface could provide "docking sites" for oxygen reduction, but it might also affect the electron transfer from Ti(III) in the negatively charged interior to the positively charged surface. Furthermore, the protonation could affect the concentration of conduction band electrons (or Ti(III)) for electron transfer at the interface. In the presence of PIM-EA-TB, the effect is further enhanced, which is likely to be linked to the pK_A of the Tröger-base amine sites in the polymer backbone at approx. pH 4.17 Protonation of the polymer backbone further enhances the rate of H2O2 production either (i) by providing protons during oxygen reduction or (ii) due to additional positive charges at the catalyst surface further enhancing electric field gradients (i.e. the rate of electron transfer). The most pronounced rate enhancement effects are observed close to pH 7 or in unbuffered media. The re-cycling of the barium titanate catalyst in PIM-EA-TB is demonstrated in Fig. S1 (ESI†).

3.2. Electron paramagnetic resonance (EPR) data for reactivity of immobilised barium titanate

EPR investigations of barium titanate provides a powerful tool to study chemical species.³⁹ Radical centres are observed for both bulk state and surface states and as a function of

treatment. First, in order to establish the bulk properties, the barium titanate powder sample was heated and dried under vacuum and then investigated at 98 K (under vacuum to remove all surface species) with/without a 300 W mercury tungsten blended UV reflector (simulating the solar spectrum) to identify bulk EPR signals (Fig. 5). The EPR response is consistent with high temperature annealed barium titanate.⁴⁰

The barium titanate powder measured at 98 K revealed the presence of Mn²⁺ trace impurities commonly found in barium titanate samples. 41 For the main signal, simulation revealed the presence of a signal exhibiting a g-tensor with axial symmetry and principal values equal to $g_{\perp} = 2.004$ and $g_{\parallel} = 1.987$. Kolodiazhnyi and Petric⁴¹ attributed a similar signal to the presence of paramagnetic lattice vacancies. A reduction of the solid can be employed to build a higher concentration of these vacancies with a temperature dependent EPR signature. 42 Interestingly, the best linewidth fit of the simulated spectrum to the experimental spectrum could only be obtained when simulating two types of Ti vacancies with the same g-tensor principal values but different g strain line broadening. This could be the reflection of Ti vacancies in two slightly different crystallographic environments. Upon 35 minutes of irradiation of the powder barium titanate with a lamp having the same emission spectrum as the solar radiation, a further signal appeared at g =1.942 (see * symbol). Kolodiazhnyi and Petric⁴¹ and the literature cited therein attributed this g value to a further Ti3+ paramagnetic centres. Next, the same barium titanate nanopowder (without heating and vacuum) was investigated (Fig. 5C). All the spectra of the impregnated paper exhibit signals corresponding to the Ti vacancies and Ti³⁺ point defects as seen previously in the simulations. A shoulder or broadening denoted "**" towards lower magnetic induction (not in the range for Ti³⁺) could be linked to surface states (associated with the oxygen species) or it could be associated with a Ti vacancy with broader line width. Both isopropanol treatment and

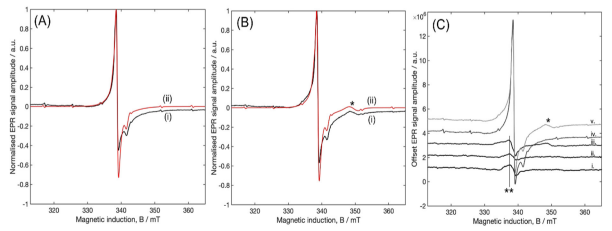


Fig. 5 (A) EPR experimental spectrum (i, black trace) and simulation (ii, red trace) of BaTiO₃ nanopowder (heated and retained under vacuum) at 98 K. (B) EPR experimental spectrum (i, black trace) and simulation (ii, red trace) of BaTiO₃ after 35 minutes of 300 W mercury tungsten blended UV reflector irradiation at 98 K. (C) EPR spectra of (i) BaTiO₃ PIM-EA-TB-embedded impregnated onto paper without solution; (ii) as before, wetted with a 30%v isopropanol aqueous solution; (iii) as before, wetted with a 30% isopropanol aqueous solution after 35 minutes of 300 W mercury tungsten blended UV reflector irradiation; (iv) $BaTiO_3$ nanopowder; (v) $BaTiO_3$ powder after 35 minutes of irradiation. All the spectra were recorded at 98 K; (iv and v were heat treated under vacuum).

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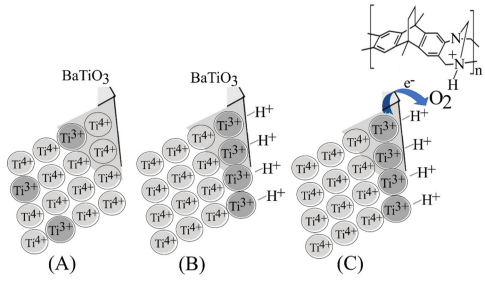


Fig. 6 Illustration of (A) the BaTiO₃ lattice, (B) the effect of protonation on double layer formation, and (C) the transition state for the oxygen reduction reaction.

exposure to light did not cause increase in the main signal (associated with the bulk).

3.3. Mechanistic model for reactivity of immobilised barium titanate

It seems likely that Ti(III) sites form at barium titanate interfaces, in particular in the presence of the reducing reagent isopropanol (eqn (1). The EPR signals did not reveal additional Ti(III) in the presence of isopropanol, but they might be dominated by bulk Ti(III) and are therefore less sensitive to surface states. The reduction of oxygen (eqn (2) then requires protons; probably involving protonation of both the barium titanate surface and the PIM-EA-TB host (see Fig. 4d). The reduction of oxygen requires two electrons (both from barium titanate) and two protons (enhanced by PIM-EA-TB protonation; Fig. 6). A similar pHdependent change in reactivity related to the polymer cage was recently proposed for nano-Pd catalysts embedded in PIM-EA-TB.²³ The presence of the additional proton in the vicinity is proposed to accelerate electron tunnelling into O2. The overall reaction (eqn (1) and (2) is pH neutral as the isopropanol oxidation can replenish both protons and electrons.

4. Conclusion

It has been shown that molecularly rigid polyamines (PIM-EA-TB) are able to host barium titanate nanoparticles for hydrogen peroxide production without capping or blocking of the catalytically active surface. Instead, an enhancement of the catalytic reaction occurs which is assigned here to a "cavity effect" inside the PIM-EA-TB that is essentially modulating the local cavity pH to switch between two states: (state 1) with the amine protonated (cavity pH < 4) the reduction of O₂ is favoured and (state 2) with the amine deprotonated (cavity pH > 4) the oxidation of iso-propanol is enhanced. Here, the reduction of O2 is likely to be the rate limiting step and

therefore the origin of the kinetic "cage" effect. There could be further kinetic effects of the PIM-EA-TB also on the catalytic decomposition of hydrogen peroxide (accompanying the formation of hydrogen peroxide), which will require further study in the future.

In the future, cavity effects in molecularly rigid host environments could be beneficial in a wider range of catalytic processes. A comparison of different polymers of intrinsic microporosity (and of other related host materials) could be useful to further explore/optimise molecular structure effects on cavity processes in catalysis. A more detailed study of similar catalytic and electrocatalytic systems will be required.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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