Demonstration of Enhanced Piezo-Catalysis for Hydrogen Generation and Water Treatment at the Ferroelectric Curie Temperature

HIGHLIGHTS
- First demonstration of the positive impact of operating near $T_c$ for piezo-catalysis
- Ultrasound applied to achieve piezo-catalysis near $T_c$
- High hydrogen production rate and degradation rate were achieved at the $T_c$
- Piezo-catalysis is a new avenue for low $T_c$ and lead-free ferroelectrics
Demonstration of Enhanced Piezo-Catalysis for Hydrogen Generation and Water Treatment at the Ferroelectric Curie Temperature

Pham Thi, Thuy Phuong,2,5,* Yan, Zhang,1,5,* Nick, Gathercole,3 Hamideh, Khanbareh,3 Nguyen Phuc, Hoang Duy,2 Xuefan, Zhou,1 Dou, Zhang,1 Kechao, Zhou,1 Steve, Dunn,4 and Chris, Bowen3,6,*

SUMMARY
Hydrogen can contribute significantly to the energy mix of the near future, as it is an attractive replacement for fossil fuels due to its high energy density and low greenhouse gas emission. A fascinating approach is to use the polarization change of a ferroelectric due to an applied stress or temperature change to achieve piezo- or pyro-catalysis for both H₂ generation and wastewater treatment. We exploit low Curie temperature (T_c) ferroelectrics for polarization-driven electrochemical reactions, where the large changes in polarization and high activity of a ferroelectric near its T_c provides a novel avenue for such materials. We present experimental evidence for enhanced water splitting and rhodamine B degradation via piezo-catalysis by ultrasonic excitation at its T_c. Such work provides an effective strategy for water splitting/treatment systems that employ low T_c ferroelectrics under the action of mechanical stress or/and thermal fluctuations.

INTRODUCTION
Hydrogen is considered as a promising and abundantly available renewable fuel with a high energy density per unit mass, with potential to act as a carbon-free energy carrier in response to the global energy crisis and need to reduce environmental pollution (Xiang et al., 2016; Zhu et al., 2018). The ability to couple piezoelectric and pyroelectric effects with electrochemical processes continues to attract attention to exploit mechanical vibrations and temperature fluctuations for hydrogen generation, water treatment, photocatalysis, and materials processing (Zhang et al., 2015, 2017; Ismail et al., 2019; Mushtaq et al., 2018). Starr et al. (Starr et al., 2012; Starr and Wang, 2013, 2015) provided a detailed overview of the fundamentals of piezo-catalysis and examined the surface electrochemistry of a piezoelectric whose polarization changed in response to a mechanical strain. To demonstrate its potential, hydrogen generation was achieved by oscillating a piezoelectric Pb(Mg_{1/3}Nb_{2/3})O_3-32PbTiO_3 (PMN-PT) cantilever in contact with deionized water (Starr and Wang, 2013). Piezoelectric materials that have been examined include zinc oxide, which have been combined with photocatalytic studies by the application of both ultrasound and light (Xue et al., 2015; Tan et al., 2015; Ma et al., 2019; Hong et al., 2016), or heat to explore pyroelectric effects due to a change in polarization with temperature (Qian et al., 2017; Tang et al., 2018). Single- and few-layer piezoelectric materials, such as MoS₂ and MoSe₂, have also been examined in detail by Wu et al. (Wu et al., 2016b, 2017, 2018c; Lin et al., 2017).

Recent work by Kakekhanı et al. (Kakekhanı and Ismail-Beigi, 2015, 2016a, 2016b; Kakekhanı et al., 2016) has focused on the use of ferroelectric materials for polarization-driven catalysis, which exhibit both piezo- and pyroelectricity. These materials are of interest because their polarization direction can be changed, thereby enabling a switching of surface chemistry. Modeling work, based on PbTiO₃, showed that splitting of water into oxygen and hydrogen is possible by thermally cycling a ferroelectric above and below its Curie temperature (T_c) when in contact with water. Initial experimental effort to demonstrate this process used pyroelectrics as an external charge source to generate a potential difference for water splitting (Zhang et al., 2019; Xie et al., 2017). However, recent work has shown that it is possible to generate hydrogen by simply placing ferroelectric particles in direct contact with water and subjecting them to cyclic heating and cooling (Belitz et al., 2017; Xu et al., 2018b; You et al., 2018a). As a model ferroelectric material, barium titanate has often been explored for tribo- (Li et al., 2019), piezo- (Belitz et al., 2017; Feng et al., 2019; Hong et al., 2010)
in relation to the results presented here therefore provide new insights into the influence of operating temperature, sono-chemical and piezo-catalytic effects, a factor that is often ignored. By careful experimental design, electric performance with respect to lead-based piezoelectric ceramics (Wu, 2018; Airimioaei et al., 2017).

Advantages include its lead-free composition, low cost, and comparable ferroelectric, piezo-, and pyro-catalytic performance in water above and below \( T_c \). Such cavitation events have also been shown to generate hydrogen from water (Islam et al., 2019), termed piezo-catalytic performance. As a result, close control of operating temperature during the sonication process (termed sono-chemical) is above the boiling point of water (\( T_b \sim 100^\circ\text{C} \)).

Although barium titanate has been widely explored for polarization-driven catalysis, its Curie temperature is above the boiling point of water (\( T_b \sim 100^\circ\text{C} \) (Harwood et al., 1947)), thereby making an evaluation of catalytic performance in water above and below \( T_b \) impossible. Other high Curie temperature ferroelectrics studied to date with \( T_c \sim 100^\circ\text{C} \) include LiNbO\(_3\) (Gutmann et al., 2012), LiTaO\(_3\) (Gutmann et al., 2012), PbZr\(_{0.75}\)Ti\(_{0.25}\)O\(_3\) (Lin et al., 2014; Feng et al., 2017), BiFeO\(_3\) (Liu and Wu, 2019; Wu et al., 2016a; You et al., 2019), NaNbO\(_3\) (You et al., 2018b; Wang et al., 2020), KNbO\(_3\) (Jia et al., 2019) (\( T_c \sim 350^\circ\text{C} \)), ZnSnO\(_3\) (Wang and Wu, 2020), and KNbO\(_3\) (Jia et al., 2019) (\( T_c \sim 434^\circ\text{C} \)) where piezo-catalytic hydrogen generation and degradation of water contaminants has been achieved at temperatures that are well below \( T_b \). One low Curie temperature material (\( T_c < 100^\circ\text{C} \)) that has been studied is barium strontium titanate (Xu et al., 2018a, 2018b), although the influence of operating temperature on piezo- or pyro-catalytic performance has yet to be examined.

In this paper we provide the first examination of the influence of working temperature, in relation to the \( T_c \), on the piezo-catalytic performance for hydrogen generation and degradation of water contaminants (rhodamine B [RhB]) under the application of ultrasound. We have selected a BaTiO\(_3\)-based solid solution based on Ba\(_{0.75}\)Sr\(_{0.25}\)TiO\(_3\) due to its relatively low Curie temperature (\( T_c \sim 42^\circ\text{C} \)), which allows us to systematically examine piezo-catalysis at temperatures (1) below \( T_c \), (2) at the \( T_c \), and (3) above \( T_c \). Additional advantages include its lead-free composition, low cost, and comparable ferroelectric, piezo-, and pyro-electric performance with respect to lead-based piezoelectric ceramics (Wu, 2018; Airimioaei et al., 2017).

RESULTS AND DISCUSSION

Although the initial pioneering work of Starr et al. (Starr and Wang, 2013) used a simple cantilever configuration to generate a strain in the piezoelectric, a number of researchers have used ultrasound to apply a mechanical deformation. A difference between these two approaches is that the application of ultrasound can lead to the formation of bubbles by cavitation of water, whose subsequent violent collapse can create localized hotspots and high pressures to generate temperatures of \( \sim 5000^\circ\text{C} \) and pressures of \( \sim 500 \text{ atm} \). Such cavitation events have also been shown to generate hydrogen from water (Islam et al., 2019), termed sono-chemical generation in this paper. As a result, close control of operating temperature during the application of ultrasound was undertaken in this work, along with the use of control samples that contained non-ferroelectric ceramic particles (termed control) or were particulate free (termed blank) to separate the sono-chemical and piezo-catalytic effects, a factor that is often ignored. By careful experimental design, the results presented here therefore provide new insights into the influence of operating temperature, in relation to the \( T_c \), on piezo-catalytic performance and demonstrates the importance of selecting an optimum operating temperature or \( T_b \) for improved water splitting or water treatment; it also demonstrates that harnessing the inherent polarization changes in ferroelectrics with stress and temperature can aid chemical conversion.

In terms of the ferroelectric Ba\(_x\)Sr\(_{1-x}\)TiO\(_3\) (BST) material used for this study, Figure 1A shows the XRD data of the synthesized powders with Miller indices indicated in the 2\( \theta \) range of 20–90\(^\circ\), where the sharp and well-defined XRD peaks indicate its high crystallinity. Only one crystalline (inset of Figure 1A) phase was detected, with no presence of secondary phase. By comparing XRD data with standard JCPDS no. 44-0093, the synthesized powders had the same characteristic peaks as tetragonal perovskite BST.
Figure 1B shows a scanning electron microscopy (SEM) image of the BST particles, where it can be seen that the particles prepared by the solid-state reaction with an equi-axed morphology exhibited a particle size of 0.2–1.2 μm. Figure 1C shows the ferroelectric polarization–electric field (P-E) hysteresis loops of a sintered BST as a function of temperature and electric field. Relative permittivity and dielectric loss of the sintered BST dense pellet as a function of temperature at a frequency of 1 kHz. (E) local ferroelectric domain switching. (a) Local hysteresis loop behavior for the amplitude and phase, (b) PFM amplitude, (c) PFM phase.

Figure 1B shows a scanning electron microscopy (SEM) image of the BST particles, where it can be seen that the particles prepared by the solid-state reaction with an equi-axed morphology exhibited a particle size of 0.2–1.2 μm. Figure 1C shows the ferroelectric polarization–electric field (P-E) hysteresis loops of a sintered BST pellet with the relative density of 98.9% at different temperatures ranging from 25 to 42°C; a sintered sample was simply used to facilitate characterization of the P-E although powders will be used for piezocatalysis. At room temperature, the saturated polarization and coercive field were 3.5 μC cm⁻² and ~1 kV cm⁻¹, respectively. With an increase of temperature, the polarization decreased and the hysteresis loops became slimmer and the hysteresis loop exhibited an almost linear response at 42°C, which can be regarded as the Curie temperature, similar to Ashish’s observation (Pradeep et al., 2010). This is also in accordance with the dielectric properties illustrated in Figure 1D, where the relative permittivity (ε) is found to initially increase with temperature, and then reaches a maximum at the Curie temperature of Tc ~ 42°C, and is then followed by a decrease in permittivity with increasing temperature that can be assigned to the transition from the asymmetric tetragonal (ferroelectric) to symmetric cubic (paraelectric) phase. The
dielectric loss (loss tangent) exhibited a similar trend but remained low in the temperature range of 26.5–65.5°C. The AC conductivity (σ) increased with frequency (f), based on the equation of $\sigma = 2\pi f\epsilon$, and there is only a limited change in AC conductivity in temperature range of 25–42°C, as shown in Figure S1. Figure 1E shows the domain pattern and the local surface displacement via piezo-force microscopy (PFM) as a function of the DC bias voltage from −10 V to 10 V. As shown in Figure 1E-a, a well-defined hysteresis response of the BST sample can be observed locally, further confirming the ferroelectricity in BST ceramic below its Curie temperature. The local surface displacement due to the converse piezoelectric effect is shown in Figure 1E-b, indicating the magnitude of the piezoelectric coefficient along the normal direction with a mean $d_{33}^*$ of ~430 pm V$^{-1}$, calculated from the PFM amplitude. An approximately 180° phase difference (from $-20^\circ$ to $-160^\circ$) can be found in Figure 1E-a, together with two different color contrasts with an ~180° phase difference, shown in Figure 1E-c, indicating the switching nature of the domains in the BST material.

In summary, the material is clearly ferroelectric and exhibits a low Curie temperature of $T_c \sim 42°C$, which makes BST a suitable candidate for the investigation of the effect of working temperature on piezoelectrically enhanced water splitting.

We now examine the influence of working temperature on the water splitting capability of BST, where the experimental setup is outlined in Figures S2 and S3. The apparatus was used to apply ultrasound to BST powder dispersed in a mixture of water and methanol, which was used as a sacrificial reagent to scavenge OH radicals (Penconi et al., 2015). As can be seen in the Figure S4, the application of ultrasound at a fixed power level and frequency (40 kHz) leads to an increase in temperature of the bulk solution with excitation time. Because sono-chemical hydrogen production can be affected by ultrasonic frequency, dissolved gas, acoustic power, and liquid temperature (Rashwan et al., 2019), they were closely controlled to ensure a stable sono-chemical hydrogen generation, as in Figure S4, to enable analysis of the effect of working temperature on piezoelectric hydrogen production. To select a specific working temperature, without altering other conditions, the hydrogen production rates at the same ultrasound on/off time ratio (1/1) with different interval times were determined using initial water bath temperatures of 23, 30, and 33°C. This ensured that the ultrasound was applied in a narrow working temperature range in relation to the $T_c$ of the material. In order to separate the amount of hydrogen that was generated by the ultrasound on and off processes, the product gases during on and off time periods were continuously swept by an argon stream and sent directly to the gas chromatography for immediate hydrogen measurement, as in Figure S2. When the ultrasound off time was greater than the retention time of the gases in the system, any hydrogen produced was fully swept out of the measurement system; when the off time is shorter than the retention time background hydrogen is detected due to accumulated hydrogen in the system. Measurement uncertainties have been estimated by at least three experiments.

Figure 2A shows the working temperature and average piezo-catalytic hydrogen generation obtained with different ultrasound on/off time ratios at different water bath temperatures. It can be seen that the degree of hydrogen evolution increases as the working temperature approaches the $T_c$ of the material; however, when the working temperature exceeds the $T_c$ the degree of hydrogen evolution decreases because the BST is no longer ferroelectric. In addition, the highest and most distinct hydrogen evolution was obtained at temperatures of 40–42°C, near the $T_c$, where the material exhibits the highest and most distinct piezoelectric coefficient. This can be explained by examination of Figure 2B (Justin Raj et al., 2011; Whatmore, 2017), which shows a schematic of the temperature dependence of spontaneous polarization of BST and its dependence with stress (piezo-coefficient). At low temperatures, below the $T_c$, the piezo-coefficients of the ferroelectric tetragonal phase are relatively small, due to the relatively small changes in polarization in response to a stress. As the temperature approaches $T_c$, greater changes in polarization are expected with an applied stress change and the piezo-coefficients increase, leading to an increase in piezo-catalytic performance due to the application of ultrasound. Above the $T_c$ the material changes to the paraelectric cubic phase and exhibits no change in polarization with ultrasound.

Figure 2C shows that use of inert $\alpha$-Al$_2$O$_3$ (as a non-ferroelectric control) did not alter the hydrogen evolution, whereas the addition of ferroelectric BST resulted in a large difference compared with both “control” ($\alpha$-Al$_2$O$_3$) and “blank” (no addition of any solid substances) tests. In addition, further evidence of no enhancement in hydrogen production at a working temperature that exceeds the $T_c$ is seen in Figure 2D, indicating that a loss in the polarization of the material (see Figures 1C and 1D) correlates with a reduction in hydrogen production. Clearly, there is a strong link between the temperature change, working temperature, and the hydrogen production in the presence of BST, whereas the control and blank test is not
influenced, confirming again the piezo-catalytic activity of BST. The dispersion of BST particles in an aqueous solution and powder microstructure morphology after ultrasound excitation are shown in Figure S5, which indicate the well-dispersed state of BST particles in the solution and no significant change in morphology due to the application of ultrasound. The fact that the efficiency of the chemical reaction was reduced when the working temperature was higher than the $T_c$ indicates that any increased thermal motion of the dispersed powder due to the increased temperature was not the primary reason for the improved catalytic performance (Mushtaq et al., 2019).

Figure 3 shows the proposed mechanism of the generation of surface charge induced by the application of ultrasound and generation of cavitation bubbles. In an ultrasonic bath, cavitation bubbles (Haar, 2015) can be formed as a result of the high-intensity ultrasound, where the cavitation bubbles expand during the rarefaction and collapse in the compression of the sound wave; see also Figure S6. A shock wave or microjet is produced from the collapsed bubbles, which leads to a high temperature and pressure (Kenneth and Suslick, 1998) with intense local heating of 5000 K, large pressures of 500 atm, heating/cooling rates of $> 10^9$ K sec$^{-1}$, and liquid jet streams of 360 km h$^{-1}$ (Suslick, 1990). However, in the presence of small collapsed water bubbles, typically 10–20 $\mu$m in size (Xu et al., 2015), the temperature of BST particles are likely to rapidly fall to the working temperature of the water bath and monitored in real time. Therefore, it is believed that the BST particles in the ultrasonic bath can be subjected to a compressive load ($\sigma$) normal or perpendicular to the polarization direction of a ferroelectric domain, which can act to change the polarization of the BST particles, and this has a greater effect near the $T_c$. It is of interest to consider the differences in the generated voltage and charge as the temperature increases up to the Curie temperature. The voltage ($V$) is determined by the piezoelectric voltage coefficient, $g_{ij} = d_{ij}/\varepsilon$, and is a measure of electric field per unit stress; the $g_{ij}$ coefficient decreases with temperature as a result of an increase in permittivity. The level of charge ($Q$) is determined by the piezoelectric charge coefficient, $d_{ij}$, which reaches a maximum at its $T_c$ (Miclea et al., 2007; Chen et al., 2014). Based on Faraday’s law, $m = QM/Fz$, where $m$ is the mass of $H_2$, $Q$ is the total
electric charge, \( F = 96,485 \, \text{C mol}^{-1} \) is the Faraday constant, \( M \) is the molar mass of the substance, and \( z \) is the valence number of ions of the substance; the amount of \( \text{H}_2 \) and RhB degradation product depend on the charge generated, whereas the threshold voltage is the driving force for the chemical reactions. The large change of polarization with stress and high \( d_33 \) near the \( T_c \) is therefore beneficial to increase the quantity of charge and \( \text{H}_2 \) generated. The overall mechanism generates a positive and negative charge on opposing surfaces (Equation 1) (Xu et al., 2018b).

\[
\text{BST} \rightarrow \text{BST} + q^- + q^+ \quad \text{(Equation 1)}
\]

The positive pyroelectric charges \( (q^+) \) are able to oxidize water molecules at the surface of the BST particle, thereby producing \( \text{H}^+ \) and \( \text{O}_2 \) (Equation 2):

\[
2q^+ + \text{H}_2\text{O} \rightarrow 2\text{H}^+ + \frac{1}{2}\text{O}_2 \quad \text{(Equation 2)}
\]

The \( \text{H}^+ \) can react with the negative pyroelectric charges \( (q^-) \) to create hydrogen (Equation 3).

\[
2\text{H}^+ + 2q^- \rightarrow \text{H}_2 \quad \text{(Equation 3)}
\]

Above the \( T_c \) the polarization is lost and no pyroelectric charges are produced (Equation 1) and the piezo-catalytic activity is therefore reduced, as in Figures 2A and 2D.

Figure 4 and Table S1 provide a summary of the hydrogen evolution rates using different methods reported in the literature. It is clear that by optimization of the working temperature in relation to the \( T_c \) a promising piezo-catalytic hydrogen production rate of 3.59 mmol/g/h is achieved, which is much larger than those obtained from a simple pyro-catalytic effect from BST (7.8 \( \mu \)mol/g/h) (Xu et al., 2018b) and from other piezoelectric materials (Wang and Wu, 2020; You et al., 2019; Su et al., 2018), with a Curie temperature of 32°C \((\text{Ba}_{0.7}\text{Sr}_{0.3}\text{TiO}_3)\) (Xu et al., 2018b), 820°C \((\text{BiFeO}_3)\) (Kiselev et al., 1963), 622°C \((\text{MoS}_2)\) (Xia et al., 2016), and >700°C \((\text{ZnSnO}_3)\) (Zhang et al., 2010). The production rate also remained stable for a testing period of up to 700min, see Figure S7.

A similar sensitivity of performance in relation to \( T_c \) is also observed in the case of sono-piezo-catalytic degradation of RhB, as can be seen in Figure 5A. The highest working temperature was observed in the case of ultrasound with a 45 s on/off interval, 41°C, respectively, exhibiting the highest RhB degradation of 58%. The degradation rate of RhB then decreased with an increase in the time interval due to decrease in working temperature. Figure 5B shows that the characteristic adsorption peak of RhB gradually decreased with increasing reaction time and complete degradation of the RhB could be achieved after 120 min. From the above results, it can be concluded that approaching \( T_c \) is the most important strategy that should be considered for maximizing the pyro-catalytic effect. Figure 5C illustrates the kinetics of the RhB degradation, where a linear relationship between \( \ln(C_0/C_t) \) and irradiation time for the degradation
of RhB can be fitted to \( \ln(\frac{C_0}{C_t}) = kt \), where \( C_0 \) and \( C_t \) are the RhB concentration at time \( t_0 \) and \( t \) respectively, and \( k \) is the apparent first-order rate constant (\( \text{min}^{-1} \)). The slope of the straight line represents the kinetic constant \( k \) with a calculated value of 0.0245 min\(^{-1}\) and \( R^2 \) of 0.9967, implying a good catalytic activity that is comparable to that of commercial Degussa P25 for sono-catalytic degradation of RhB (Zhou et al., 2015).

Conclusion
In summary we have provided the first experimental demonstration of the importance of working temperature, in relation to the Curie temperature for piezo-catalytic hydrogen generation and degradation of water contaminants (RhB) under ultrasound. The piezo-catalytic performance of the ferroelectric materials increases significantly near the \( T_C \) due to an increase in piezo-coefficients and the impact of cavitation of the local stress; further work to examine any potential pyroelectric contributions due to the rapid localized temperature changes at cavitation sites would be of interest. A wide variety of lead-free ferroelectrics are currently under investigation (Zheng et al., 2018), and although the low \( T_C \) of particular systems makes them inappropriate for conventional sensor or actuator applications, they would be of interest for such piezo-catalysis. Under optimum conditions of operation near \( T_C \) the addition methanol at only 4% can enhance the hydrogen evolution with negligible generation of other impurities. Consequently, a high hydrogen production rate of 10.83 \( \mu \text{mol/h} \) was achieved, which is much higher than previously reported, such as 3.4 \( \mu \text{mol/h} \) by Wang et al. (Wang et al., 2010) (see Table S1). We also demonstrate the need for careful control of experiments to separate both sono-chemical and pyro-chemical effects. It is also of interest to explore other modes of operation such as piezo-photo-catalytic effects (Tan et al., 2015; Huang et al., 2017) near \( T_C \) or mixing ferroelectric particles of different \( T_C \) to enable operation over a range of temperatures. Of particular interest here is that for the materials selected for this study, \( \text{Ba}_x\text{Sr}_{1-x}\text{TiO}_3 \) the Curie temperature can be tailored from \(-200^\circ\text{C}\) to \(120^\circ\text{C} \) (Airimioaei et al., 2017; Mbarki et al., 2014) and provides ample opportunities to tune the Curie temperature for a range of working temperatures for optimum piezo- and pyro-catalysis.

Limitations of the Study
Particles of ferroelectric were suspended in water in this work, and methods to disperse larger amounts of particulates in water are needed to improve the amount hydrogen formed or allow scale-up for water treatment. Potential avenues of research could include porous materials. Although ultrasound was used in this work, harvesting ambient sources of energy would be beneficial; this could include inducing cavitation from water flow or using temperature fluctuations.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.
DATA AND CODE AVAILABILITY
The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101095.

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AUTHOR CONTRIBUTIONS
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DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Demonstration of Enhanced Piezo-Catalysis
for Hydrogen Generation and Water Treatment
at the Ferroelectric Curie Temperature

Pham Thi, Thuy Phuong, Yan, Zhang, Nick, Gathercole, Hamideh, Khanbareh, Nguyen Phuc, Hoang Duy, Xuefan, Zhou, Dou, Zhang, Kechao, Zhou, Steve, Dunn, and Chris, Bowen
Supporting Information

Transparent Methods

Material preparation:
Ba_{0.75}Sr_{0.25}TiO_3 (BST) powders were prepared by a solid-state reaction. Analytical grade (Sigma Aldrich) barium carbonate (BaCO_3, 99%), strontium carbonate (SrCO_3, 99%), and titanium dioxide (TiO_2, 99.9%) were selected as starting materials and weighed according to their stoichiometric ratio. The above mixtures were calcined at 1200 °C for 3 h, followed by additional ball-milling for 24 h. The milled powders were mixed with 1 wt.% poly(vinyl alcohol) (PVA) binder and dried in an oven at 60 °C. To form dense materials for characterisation, the powders were uni-axially cold-compacted to form pellets of 10 mm in diameter and 1 mm in thickness. The pellets were first heated to 500 °C for 3h to remove the binder and then sintered at 1325 °C for 4 h.

Characterization:
The phase structure of the BST powders was examined by X-ray diffractometer (BRUKER D8-Advance, USA) with Cu radiation with 2θ ranging from 20°-70°. The morphology of the BST powders was examined by a scanning electron microscopy (SEM, JSM6480LV, Tokyo, Japan). For electro-physical measurements of sintered materials, silver paint was coated on both working faces of the sintered samples to form electrodes. The temperature dependence of the dielectric constant (relative permittivity) and polarization of the unpoled pellets were studied in a temperature range of 26-67 °C using an impedance analyzer (Solartron 1260, Hampshire, UK) at a frequency of 1 kHz and a Radiant RT66B-HVi Ferroelectric Test system at hysteresis period of 10 ms, respectively. Piezo force microscopy (PFM) in contact mode of an atomic force microscope (NanoManTM VS) with a conductive Pt/Ir-coated Si cantilever (SCM-PIT) was used to investigate the piezo-response phase and amplitude of BST sample. The ceramic sample was polished, thermally etched, and ground to 0.2 mm before measurement. A direct current (DC)
voltage from -10 to 10V was applied superimposed on an AC modulation voltage during polarisation switching.

**Hydrogen production experiments:**
A mass of 1 mg of BST powder was added and dispersed into a 20 mL headspace vial containing 10 mL of distilled water and methanol (Merk) mixture. After sealing tightly, it was placed in the center of an ultrasonic cleaner bath (60W-TP01-Taiwan Total Meter) and purged with argon for 15 min to completely remove air prior to being sonicated for a desired reaction time at a frequency of 40 kHz (see Figure S2). For online measurement, the generated hydrogen was continuously swept by argon stream at about 12 ml/min and passed to a 120 mL condensation bottle before sending to a gas chromatograph (HP 5890 Series II, Agilent) equipped with a thermal conductivity detector and RT-Msieve 13X capillary column (30 m x 0.32 mm, Thames Restek) by ChemStation Software (see Figure S3). Actual flow rates were verified with a soap-film bubble flowmeter (Hewlett-Packard) prior to each run. The hydrogen production rate was calculated using a calibration curve that was made by using different concentrations of hydrogen diluted in argon. During the experiment, the ultrasonic bath temperature was controlled by circulating cooled water back to the ultrasonic bath. The control (non-ferroelectric particulate addition) tests were performed at the same conditions with 1 mg of α-Al₂O₃ (Merk) as an inert material and blank tests without adding any solid particle to the reaction mixture. This allowed the *sono-chemical* contribution to hydrogen generation be determined. The uncertainties have been estimated by at least three experiments. The *piezo-catalytic* hydrogen generation was calculated by simple subtraction of hydrogen evolution rates obtained from experiments with BST and blank tests due to no statistical difference in hydrogen generated from control and blank tests, then divided by the amount of BST.

**Rhodamine B degradation experiments:**
A mass of 2 mg of BST powder was added and dispersed into a 20 mL headspace vial containing 10 mL of RhB solution (10 mg/L), which has been widely chosen as a model contaminant based on its popularity among research groups (Rochkind et al., 2015). By conducting mutually exclusive experiments without
applying ultrasound, it was found that the direct photolytic degradation, adsorption and thermal
degradation of RhB on BST are negligible. Thus, all further experiments were performed at normal
light condition and without adsorption pre-treatment. The absorption spectra of the treated samples
were acquired with a UV-Vis spectrophotometer.

![Figure S1 AC conductivity of BST with different frequencies and temperatures, related to Figure 1.](image-url)
Figure S2. Schematic image of the experimental setup. 1 – Ultrasonic bath; 2 – Sample vial; 3 – Inside temperature sensor; 4 – Water bath temperature sensor; 5 – Timers; 6 – Peristaltic pump; 7 – Cooling coil; 8 – Cooling bath temperature sensor; 9 – Interface; 10 – Computer; 11 – Flexible stand, related to Figure 2.
Figure S3. (A) Online hydrogen measurement setup. (B) Representative GC-TCD chromatogram of the produced sample at optimum conditions. 12 – Three ways valve; 13 – Condensation bottle, related to Figure 2.

Figure S4. Hydrogen generation for blank test, with sono-chemical produced hydrogen shown to be temperature independent, related to Figure 2.
Figure S5. Images of the aqueous solution before and after powder addition. (A) pure water, (B) BST aqueous suspension after the application of ultrasound excitation, inset shows no precipitation can be seen on the bottom side, (C) SEM image of the BST powder after the application of ultrasound, related to Figure 2.

Figure S6. Schematic of ultrasound applied to ferroelectric particle, related to Figure 3.
Table S1 Rate of hydrogen evolution under different conditions, related to Figure 4.

| Effect                  | Material      | Hydrogen evolution rate | Reference               |
|-------------------------|---------------|-------------------------|-------------------------|
|                         |               | µmol/h | mmol/h/g |                         |
| Pyro-catalytic          | BST           | n/a     | 0.0078\(^a\) | (Xu et al., 2018)       |
| Sono-chemical           |               | 3.4\(^b\) | n/a      | (Wang et al., 2010)     |
|                         |               | 10.83 ± 0.13\(^c\) | n/a      | This work               |
| Sono-catalytic          | Au/TiO\(_2\)  | 282.3\(^b\) | 3.8\(^b\) | (Wang et al., 2010)     |
| Piezo-photo-catalytic   | KNbO\(_3\)/MoS\(_2\) | n/a     | 0.096\(^d\) | (Jia et al., 2019)      |
| Sono-piezo-catalytic    | BST           | 14.42 ± 0.15\(^c\) | 14.42 ± 0.15\(^c\) | This work               |
|                         | ZnSnO\(_3\)   | n/a     | 3.5\(^e\)  | (Wang and Wu, 2019)     |
|                         | MoS\(_2\)     | n/a     | 0.028\(^f\) | (Su et al., 2018)       |
|                         | BiFeO\(_3\)   | n/a     | 0.026\(^g\) | (You et al., 2019)      |
| Extracted piezo-catalytic effect | BST | 3.59\(^c\) | 3.59\(^c\) | This work               |

\(n/a\) abbreviates either not available or not applicable

\(^a\) Thermal cycle is 10 min with thermal fluctuation of 25-50 °C, concentration of methanol is 20 vol%

\(^b\) Concentration of methanol was 4 vol%, ultrasound source of 40kHz at 50 W

\(^c\) Concentration of methanol was 4 vol%, ultrasound source of 40kHz at 60 W, ultrasound on/off time ratio is 15/15s with thermal fluctuation of 40.5 - 42 °C

\(^d\) Triethanolamine was used as a sacrificial reagent (15%), ultrasound source of 40kHz at 110 W

\(^e\) Anhydrous alcohol was used as a sacrificial reagent (50 vol%), ultrasound source of 40kHz at 250 W

\(^f\) Fe\(^{2+}\) was used as inorganic oxidizable sacrificial agent, ultrasound source of 40kHz at 140 W

\(^g\) Na\(_2\)SO\(_3\) was used as a sacrificial reagent (0.05M), ultrasound source of 45kHz at 50 W
Figure S7. Piezo-catalytic hydrogen generation with time on stream, related to Figure 4.

In this experiment, a condensable bottle (see Fig. S2-A) was used to determine a stable hydrogen production rate. Thus, there is a spike at ~100 mins which is the required duration time for the H₂ to fully fill the bottle. H₂ production was found to be affected by methanol concentration, therefore a subsequent decrease (200min) may be a result of a decrease in methanol concentration due to its vaporisation. At long times, >200min the sample volume is likely to decrease during ultrasound excitation, resulting an increase in inside temperature (outside temperature and ultrasound power were constant), thus, an increase of H₂ production rate may occur for longer durations.
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