High-Pressure Phase Transitions of Morphologically Distinct Zn$_2$SnO$_4$ Nanostructures

Partha Pratim Das, P. Sujatha Devi, Douglas A. Blom, Thomas Vogt, and Yongjae Lee

1Department of Earth System Sciences, Yonsei University, Seoul 120749, Korea
2Sensor and Actuator Division, CSIR-Central Glass and Ceramic Research Institute, Kolkata 700032, India
3NanoCenter & Department of Chemical Engineering, and NanoCenter & Department of Chemistry & Biochemistry, University of South Carolina, Columbia, South Carolina 29208, United States
4Center for High Pressure Science and Technology Advanced Research, Shanghai 201203, China

ABSTRACT: Many aspects of nanostructured materials at high pressures are still unexplored. We present here, high-pressure structural behavior of two Zn$_2$SnO$_4$ nanomaterials with inverse spinel type, one a particle with size of $\sim$7 nm [zero dimensional (0-D)] and the other with a chain-like [one dimensional (1-D)] morphology. We performed in situ micro-Raman and synchrotron X-ray diffraction measurements and observed that the cation disordering of the 0-D nanoparticle is preserved up to $\sim$40 GPa, suppressing the reported martensitic phase transformation. On the other hand, an irreversible phase transition is observed from the 1-D nanomaterial into a new and dense high-pressure orthorhombic CaFe$_2$O$_4$-type structure at $\sim$40 GPa. The pressure-treated 0-D and 1-D nanomaterials have distinct diffuse reflectance and emission properties. In particular, a heterojunction between the inverse spinel and quenchable orthorhombic phases allows the use of 1-D Zn$_2$SnO$_4$ nanomaterials as efficient photocatalysts as shown by the degradation of the textile pollutant methylene blue.

INTRODUCTION

The distinct behaviors of nanostructured materials from those of their bulk counterparts are the prime motivation behind research on nanomaterials in various fields. The interplay among particle sizes of the building blocks in the nanoregime, the different atomic structures, and the surface and interfacial properties of the grains are the deciding factors in the distinctive functional behavior of nanomaterials. In the pursuit of new phenomena in nanomaterials, high pressure has been employed recently to modify the atomic arrangements and interactions of materials. Understanding the pressure-induced structural stability of various types of nanomaterials and the concomitant tuning of their physicochemical properties is important from both a fundamental and application point of view.

Over the past few decades, semiconducting nanomaterials of different types and structures have greatly contributed to significant progress in nanoscience and technology due to their salient and flexible opto-electronic, photonic, magnetic, and mechanical features. As far as the exploration of semiconducting nanomaterials under high pressure is concerned, studies on a variety of materials, namely, the group (IV) elements C, Si, and their compound SiC; group (II−VI) compounds such as ZnS, ZnSe, CdS, CdSe, and CdTe; group (IV−VI) PbS; group (III−V) GaN and AlN; the superhard material, B$_4$C$_2$N$_2$ binary oxides such as TiO$_2$, ZnO, SnO$_2$, Fe$_2$O$_3$, and CeO$_2$; the rare-earth oxide, Ho$_2$O$_3$; wide band-gap oxides such as $\beta$-Ga$_2$O$_3$ and Y$_2$O$_3$; p-type compounds including CuO, CoO, and MnS; n-type BaTiO$_3$; and narrow band-gap layered group (V−VI) semiconductors such as Bi$_2$Te$_3$, have been performed.

Most of these studies have shed light on the interesting kinetics of pressure-induced first-order, solid−solid structural transformations, compressibilities, bulk moduli, and stiffness or hardness of the materials. The thermodynamics of phase transformations and relative stabilities of the phases have also been noted in several studies. Although there are conflicting trends in the reported transition pressures relating to the Hall−Petch effect that is found as bulk materials are reduced to smaller crystallites, a significant influence from nanosized particles or grains has been commonly suggested as the cause for the dissimilar types of nucleation, growth dynamics, phase transition pathways, and even sequences of the phase transitions or amorphizations of semiconducting materials under high pressure. A specific size, at which the typical nanoscale effects start to occur in materials, has also been defined as their respective “critical size” in several cases. The contributions of the nanoscale-induced differences in the surface energies of the relevant phases mainly account for the stabilities of the corresponding structures. In this context, the impacts of various microstructural features, for instance, the distinct shape or...
morphology, dimension, and homogeneity, of the materials are often found to coincide with the size effects of the nanocrystals in many studies.31,44,46,48–51 The microstructure-induced strains appearing at the contact points of the grains of the materials may cause significant structural distortions, which also contribute to the transition pressure and phase stability.14,49

In the present study, the simultaneous effects of the size and microstructure have been investigated on the high-pressure phase transition of a specially designed ZnO–SnO2-based multi-cation oxide, Zn2SnO4. The wide band gap, n-type semiconducting Zn2SnO4 is highly acclaimed as a potential candidate for various applications, such as dye-sensitized solar cells, transparent conducting oxides, photocatalytic degradation, humidity and combustible gas sensing, and Li-ion batteries.40–42 Based on their work, the pressure-induced phase transition of spinel Zn2SnO4 to orthorhombic phases was first experimentally detected at 10 GPa, where remarkably, no structural transition was observed.33 The ternary oxide, Zn5SnO4Fe, which has cubic (Fd3m) symmetry, belongs to the (A2+)[B4+A2+]O4-type inverse spinel oxides in the bulk state, where half of the A cations populate the tetrahedral sites and the other half occupy the octahedral sites along with all of the B cations.46 Several spinel oxides possess complicated disordering phenomena involving the redistribution of cations A and B over the tetrahedrally and octahedrally coordinated sites. There have been reports claiming a significant role of this cation reordering in spinel oxides with regard to the structural stability and diffusion or diffusionless phase transformations at high temperatures.47–49 Few studies have highlighted a similar phenomenon for several bulk spinel oxides at high pressure.47–49 The Zn2SnO4 nanoparticles with grain sizes of ~7 nm used in this present study reveal cation disorder. Therefore, this represents a unique occasion to characterize the phase transformations associated with cation reordering of a nanosized inverse spinel oxide, which is another important aspect of the present study.

### RESULTS AND DISCUSSION

**Microstructural and Optical Characterizations of the Samples under Ambient Conditions.** A bright field transmission electron microscopy (TEM) image of the hydrothermally synthesized Zn2SnO4 sample dispersed in ethanol is presented in Figure 1a. The high-resolution image shows the monodispersed spherical particles with sizes of less than 10 nm in all dimensions. The particles are marked with yellow dotted lines for a clear demarcation. At the same time, the 0-D nanoparticles are found to be aggregated in the final Zn2SnO4 sample studied here as revealed by the high-angle annular dark-field-scanning TEM (HAADF-STEM) images in Figure 1b. As a consequence, a moderate surface area of...
∼36.71 m²/g was obtained for the Zn₂SnO₄ nanoparticles prepared by the hydrothermal method from the Brunauer–Emmett–Teller (BET) analysis. In addition, the characteristic type-III isotherm that is observed in the N₂ adsorption–desorption studies (Figure 1c) along with an insignificant pore size distribution (PSD) as calculated by the Barrett–Joyner–Halenda (BJH) method (Figure 1c, inset) further support the nonporous nature and compactness of the nanoparticles of the Zn₂SnO₄ sample.

The Zn₂SnO₄ sample prepared from the temperature-assisted solid-state reaction between ZnO and SnO₂ appears to have a self-assembled elongated chain-like unique morphology. The STEM image of a small unit of the typical chain-like particles of Zn₂SnO₄ and the TEM image that includes several similar particles assembled together are displayed in Figure 1d,e, respectively. The actual width of the individual chain-like particles measured considering only their nonoverlapping sections varies over the range of 50−100 nm (marked with yellow double arrows). The dark contrast in both the STEM and TEM images of the 1-D particles indicates that the Zn₂SnO₄ nanostructures formed by the solid-state method are highly dense in nature and have no Kirkendall pores. This, again, is supported by the type-III N₂ adsorption–desorption isotherm exhibited by the dense Zn₂SnO₄ nanostructures (Figure 1f). The BET surface area obtained from the sample was found to be very low at ∼4.23 m²/g. However, the BJH PSD, shown in the inset of Figure 1f, contrarily reveals the presence of pores with a wide distribution of sizes varying from ∼5 to ∼60 nm even in these dense Zn₂SnO₄ nanostructures. The intergrain voids (marked with red arrows) among the chain-like particles assembled with each other, and that assembly should be the only source of introducing meso- to microporosity in the sample (Figure 1e), which makes this particular morphology more exceptional for studying at high pressure.

The order–disorder phenomena in the local structures of the samples were verified using HAADF-STEM real-space imaging. A high-resolution HAADF-STEM image from the 1-D Zn₂SnO₄ nanostructures viewed along the (100) direction is displayed in Figure 2a. The inverse-spinel crystal structure of Zn₅SnO₆ (x, y, z = 1 unit) that is superimposed on the experimental HAADF-STEM image matches with the Z-sensitive contrast of the Zn and Sn columns that define the ordered distribution of Zn²⁺ and Sn⁴⁺ over the lattice sites (Figure 2a, inset). The sites with brighter contrast in the image, marked with yellow balls, correspond to the Sn atoms, which have a higher atomic number (Z) compared to Zn, and the sites of the Zn atoms are marked with red balls. In contrast, the HAADF-STEM image of the 0-D nanoparticles showed significantly lower contrast. This interesting phenomenon suggests a lack of ordering between the Zn and Sn sites in the hydrothermally synthesized Zn₂SnO₄ nanoparticles. The possibility of a redistribution of Zn²⁺ and Sn⁴⁺ in the octahedral and tetrahedral crystal sites of the inverse spinel Zn₂SnO₄ is considered, in this regard, as an effect of the reduction of particle size because we have <10 nm particle sizes. The imperfect alignment of the Zn and Sn columns as related to the aforementioned disorder may lead to the low contrast observed in the HAADF image. The distorted local structure of this particular 0-D nanosized Zn₂SnO₄ was further confirmed from its unusual Raman phonon A₁g(2) mode at ∼626 cm⁻¹, which is the characteristic vibrational mode for a nonequilibrium cation distribution in the inverse spinel structure (Figure 2c). The other Raman active bands seen in Figure 2c can be attributed to the A₁g(1), F₂g(3), F₂g(2), E₁g, and F₂g(1) symmetries of the typical inverse spinel...
structure of Zn$_2$SnO$_4$ and appeared at similar positions for both samples. The measured ratio of the intensity of the F$_{2g}(2)$ and F$_{2g}(3)$ bands of the 1-D nanoparticles with peaks at $\sim$529 and 559 cm$^{-1}$ is 1.3:1. However, the same with an inverse intensity order (F$_{2g}(2)/$F$_{2g}(3) = 1.042$) indicates the possibility of higher [B] site populations of Zn$^{2+}$ instead of Sn$^{4+}$ in the 0-D Zn$_2$SnO$_4$ nanoparticles. Moreover, the exceptionally broad Raman spectral lines further suggest that the 0-D Zn$_2$SnO$_4$ nanoparticle has a deformed MeO$_6$ octahedra as well as MeO$_4$ tetrahedra (Me = Zn or Sn). To eliminate the likelihood of any chemical disorder in this case, the chemical compositions of the samples were compared with a reported mixture of ZnO and SnO$_2$ at a ratio of 2:1, based on the X-ray photoelectron spectroscopy (XPS) analysis. It is worth mentioning that the Zn 2p$_{3/2}$ spectra of both Zn$_2$SnO$_4$ samples studied here are found to be nearly identical to that of the standard mixture with respect to the positions of their binding energies ($\sim$1020.92, $\sim$1020.91, and $\sim$1021.4 eV for the 0-D Zn$_2$SnO$_4$ nanoparticles, 1-D nanostructures, and the standard mixture, respectively) and symmetric spectral shapes, as represented in the inset of Figure 2c. This elucidates that the samples, even though synthesized by different methods, are chemically homogeneous with an atomic ratio of 2 for Zn to Sn. The XPS investigation also confirms the influence of no chemical disorder on the nonuniform structure of the 0-D Zn$_2$SnO$_4$ nanoparticles.

Therefore, based on the HAADF, Raman, and XPS analyses, it can be stated that, while the 1-D Zn$_2$SnO$_4$ nanostructures have the inverse spinel structure, the cation distribution in the 0-D nanoparticles have a partially inverse spinel structure with the chemical formula $(\text{Zn}_n\text{Sn}_{1-n})^{2+}[\text{Sn}_m\text{Zn}_{2-m}]^{2+}\text{O}_4$, where the value of the inversion parameter, “$\delta$” is between 0 and 1. A decrease in the optical band gap from $\sim$3.85 to $\sim$3.51 eV was estimated from the redshifted absorption edge in the diffuse reflectance (DR) of the 0-D nanoparticles compared to that of the 1-D nanostructures yet having the larger particle sizes, as exhibited in the inset of Figure 2d. This observation corroborates the first principles study performed by Wei et al. on the anomalous band gap reduction as a function of the cation inversion for Sn$^{4+}$/O$_4$-type spinel oxides, such as Zn$_2$SnO$_4$. The emission properties of the as-prepared samples were measured thereafter (Figure 2d, inset). The emission maxima appeared at $\sim$328 nm for the 1-D nanostructures, whereas that for the 0-D nanoparticles shifted to the mid-UVA region at $\sim$358 nm. The observed emissions of both Zn$_2$SnO$_4$ samples refer to their intraband electron transitions only.

0-D Zn$_2$SnO$_4$ Nanoparticles under High Pressure: Effects of Size. The pressure-induced structural behavior of the 0-D nanosized Zn$_2$SnO$_4$ accompanying the nonequilibrium structure was investigated by in situ Raman and X-ray diffraction (XRD) studies. Based on earlier reports, we anticipated an order—disorder-mediated structural transformation in these studies. Therefore, the variation in the relative intensities of the A$_{1g}(2)$ (red shaded area, Figure 3a) and F$_{2g}(2)$ modes versus that of the F$_{2g}(3)$ mode was monitored at elevated pressures as shown in Figure 3b. The impact of high pressure on cation reordering was negligible because there is no significant pressure-dependent change in the intensity of the A$_{1g}(2)$ peak. The F$_{2g}(2)/$F$_{2g}(3)$ ratio remained almost constant as well, up to $\sim$40 GPa, after a minute initial enhancement at $\sim$8–10 GPa. The F$_{2g}(2)$ and F$_{2g}(3)$ peaks obtained by multipeak fitting of the Raman spectrum at ambient, $\sim$10 and $\sim$37 GPa are displayed in the inset of Figure 3b. An indisputable stress arises to make more compact contact between the spherical nanograins and may have induced some disorder at lower pressure (also discussed later). However, unperturbed structural disorder under high pressure usually leads to a martensitic phase transformation through a diffusionless mechanism, which should promote the possibility of a faster phase transition in spinel structures, such as Zn$_2$SnO$_4$. In contrast, no salient modification in the Raman modes indicating a new phase are observed for the sample up to $\sim$40 GPa, except for a regular shift of the bands toward higher frequencies (Figure 3a). This exceptional behavior of the 0-D Zn$_2$SnO$_4$ nanoparticles is further
confirmed by the XRD patterns shown in Figure 3c, which exhibit only a systematic pressure-induced shift in the 2θ values toward higher angles along with certain peak broadening. A significantly high bulk modulus (Bo) of ~241.5(2) GPa can be determined, thereafter, using the second-order Birch–Murnaghan equation of state, where Kr is fixed at 4 (Figure 3d). Therefore, the 0-D small ~7 nm particles are stable and do not undergo a martensitic transformation. This study introduces diffusionless cation reordering as an important size effect during the pressure-induced phase transformation of this spinel oxide. Further studies are underway to determine whether there is any “critical size” for the Zn2SnO4 nanoparticles, where the cation disorder in the structure persists, while at the same time influencing either the diffusional or diffusionless phase transformation at higher pressures.

1-D Zn2SnO4 Nanostructures under High Pressure: Effects of Morphology. The pressure-dependent structural transformation of the 1-D Zn2SnO4 nanostructures with an elongated chain-like morphology was explored using the same pressure range of up to ~40 GPa. Interestingly, two successive modifications of the inverse spinel structure of the sample at ambient conditions (space group: Fd3m; calculated lattice parameter: a = 8.66(1) Å) can be seen in the XRD patterns as a function of pressure, shown in Figure 4A. (The small impurities of ZnO (square shaped mark) and SnO2 (round shaped mark) are not considered in the analysis.)

As a splitting of the diffraction peak corresponding to the (400) plane into two peaks was found to begin at ~10 GPa, which can be attributed to a tetragonal distortion of the cubic inverse spinel structure. The lattice parameters calculated by whole-profile fitting of the corresponding XRD pattern are a = 8.58(3) Å and a = b = 6.09(1) Å, c = 8.50(1) Å for the inverse spinel and tetragonal structures, respectively. The distortion became stronger with increasing pressure up to ~26 GPa. The analogous distortion, albeit ambiguously characterized as an intermediate phase by Shen et al., was followed by a further phase transition to the ambient-pressure structure of orthorhombic CaFe2O4 at ~32 GPa for the Zn2SnO4 nanowires with an average diameter of ~150 nm. Similarly, the onset of a second phase transition can also be observed at ~30 GPa for the chain-like Zn2SnO4 with an average width of ~100 nm, indicating a comparable surface area-to-volume ratio to that of the Zn2SnO4 nanowires studied by Shen et al. However, it is worth noting that a new denser high-pressure orthorhombic CaFe2O4 structure appeared, which is equivalent to the CaFe2O4 structure that occurred at ~63 GPa, as reported by Yamanaka et al. The analysis by Yamanaka et al. provides the same Pnnm space group symmetry but with a three-fold larger c-axis dimension (c = 8.533(1) Å) for this new phase compared to that of the orthorhombic CaFe2O4 structures that appear at lower pressures. For instance, the calculated values for “c” are 2.889(3) and 3.019(2) Å at ~42 and 0.0001 GPa, respectively. An identical “c”, 8.448(6) Å corresponding to the new phase, was obtained in our case for the chain-like Zn2SnO4 nanostructures by whole-profile fitting analysis of the XRD pattern at ~30 GPa. The complete transformation into the new phase (c = 8.347(2)) is found to take place at ~38 GPa though the gradual disappearance of the tetragonal structure at ~32 GPa followed by the inverse spinel structure at ~34 GPa. The high pressure phase of the orthorhombic CaFe2O4-type structure is quenchable and exhibits an irreversible phase transition even after releasing the pressure. The normalized unit cell volumes of the consecutive phases are shown in Figure 4C.

Finally, the selective XRD patterns mentioned above are represented in Figure S1a–d. To the extent of our knowledge, this is the first observation of the transformation of the inverse spinel structure of Zn2SnO4 into a much denser high-pressure CaFe2O4-type structure.

The pressure-induced phase transformations observed from the XRD analysis of the elongated chain-like Zn2SnO4 nanostructures were further verified by in situ Raman spectroscopic study (Figure 4B). Primarily, a substantial shift can be observed for the five first-order Raman active modes, A1g(1), F2g(3), F2g(2), E′ and F2g(1), representing the cubic

Figure 4. (A,B) Pressure-dependent XRD and Raman spectra of the 1-D Zn2SnO4 nanostructures, (C) variations in V/V0 of the evolved phases with pressure, (D) Raman spectra of the 1-D Zn2SnO4 nanostructures during decompression and (E) comparison of the calculated strains for (a) 0-D Zn2SnO4 nanoparticles and (b) 1-D Zn2SnO4 nanostructures.
Similar to the XRD results, the bands correspond to the tetragonal structure (Figure 4D). However, it has not escaped our notice that the structure highlights the involvement of a high amount of strain during the pressure-induced phase transformation of the 1-D Zn$_2$SnO$_4$ nanostructures, as reported by Gracia et al.\textsuperscript{44} However, an additional absorption feature at $\sim$790 cm$^{-1}$ can be assigned to the respective band edge of the DR spectrum and, accordingly, in the emission spectra of the recovered 1-D Zn$_2$SnO$_4$ nanostructures along with the inverse spinel phase, which results in a higher band gap of $\sim$4.19 eV compared to the as-prepared sample (3.85 eV). The low energy absorption in this case can be unequivocally considered to originate from the orthorhombic phase. Interestingly, the broad emission from the UV to visible region corresponding to this particular sample exhibits three maxima. An intense peak at $\sim$311 nm followed by a broad peak at $\sim$371 nm can be assigned to the respective band edge emissions of the inverse spinel and orthorhombic phases. Moreover, a broad visible band centered at $\sim$556 nm may originate from the strain-induced disorder in the chain-like Zn$_2$SnO$_4$ nanostructures, as observed in the Raman spectrum.

First, formation of the high-pressure orthorhombic CaFe$_2$O$_4$-type structure, yet having a comparable cohesive energy to that in the previous study, followed by disordering in the structure highlights the involvement of a high amount of strain during the pressure-induced phase transformation of the 1-D Zn$_2$SnO$_4$ nanostructures. The typical chain-like grains, which have several uneven inter grain voids (as depicted from the TEM image and BET study in Figure 1e,f), may induce extreme stress at the contact points among the grains.\textsuperscript{8,14} The inhomogeneous distribution of the stress with increasing pressure and the consequent microstrain developed at the grain boundaries are considered to play a crucial role in the phase transformation pathway of the Zn$_2$SnO$_4$ nanostructures.\textsuperscript{15} In this regard, the pressure-induced evolution of the microscopic strain based on the major (311) diffraction peak of the inverse spinel phase is also determined for the 1-D Zn$_2$SnO$_4$ nanostructures using the Stokes–Wilson equation\textsuperscript{17} and is compared with that of the 0-D nanoparticles having a nonporous and complete distribution of spherical particles mentioned earlier (Figure 4E). After an initial densification, the strain remained almost homogeneous for the 0-D Zn$_2$SnO$_4$ nanoparticles. In contrast, a much higher strain, which constantly increased with pressure and reached a maximum value at the onset pressure of the orthorhombic CaFe$_2$O$_4$ phase at $\sim$28 GPa, can be seen for the Zn$_2$SnO$_4$ sample having a chain-like morphology. A similar trend in the microstrain was also observed based on the (111), (220), (511), and (440) diffraction planes, as represented in Figure S2. The notable jump in the microstrain at $\sim$10 GPa (Figures 4E and S2) may be attributed to grain refinement of the chain-like Zn$_2$SnO$_4$ nanostructures, similar to that reported by Zhang et al.\textsuperscript{10}

Optical Characteristics of the Recovered Samples: Comparison of Their Photocatalytic Activities. The pressure-dependent modifications of the Zn$_2$SnO$_4$ samples were further isolated by successive verification of their band gaps and emission properties after the pressure was released. Only a small blueshift of $\sim$10 nm was found in the absorption edge of the DR spectrum and, accordingly, in the emission maxima of the recovered 0-D Zn$_2$SnO$_4$ nanoparticles (Figure 5A) compared to those of the ambient sample. This result suggests an extensive effect of the pressure-induced increase in the direct band gap of the spinel Zn$_2$SnO$_4$ structures, as reported by Gracia et al.\textsuperscript{44} However, an additional absorption edge of $\sim$368 nm with a much lower band gap of $\sim$3.37 eV was observed for the recovered 1-D Zn$_2$SnO$_4$ nanostructures along with the inverse spinel phase, which results in a higher band gap of $\sim$4.19 eV compared to the as-prepared sample ($\sim$3.85 eV). The low energy absorption in this case can be unequivocally considered to originate from the orthorhombic phase. Interestingly, the broad emission from the UV to visible region corresponding to this particular sample exhibits three maxima. An intense peak at $\sim$311 nm followed by a broad peak at $\sim$371 nm can be assigned to the respective band edge emissions of the inverse spinel and orthorhombic phases. Moreover, a broad visible band centered at $\sim$556 nm may originate from the strain-induced disorder in the chain-like Zn$_2$SnO$_4$ nanostructures, as observed in the Raman spectrum. The as-prepared Zn$_2$SnO$_4$ samples were compared with their respective recovered counterparts in regard to their efficiencies as photocatalysts toward the degradation of a model organic pollutant, MB, under UV illumination (365 nm, 8 W). The comparative degradation profiles of the characteristic absorption at 665 nm of MB over time are shown in Figure 5B, where $C_0$ and $C$ indicate the concentrations of MB before and after UV exposure, respectively. The rate of photodegradation efficiency is significantly altered after 35 min of UV irradiation in the order 1-D Zn$_2$SnO$_4$ nanostructures$_{\text{as prepared}}$ ($E_g \approx 3.85$ eV) $< 0$-D Zn$_2$SnO$_4$ nanoparticles$_{\text{recovered}}$ ($E_g \approx 3.62$ eV) $< 0$-D Zn$_2$SnO$_4$ nanoparticles$_{\text{as prepared}}$ ($E_g \approx 3.51$ eV) $< 1$-D Zn$_2$SnO$_4$ nanostructures$_{\text{recovered}}$ ($E_g$ inverse spinel). $E_g$ values for the different nanostructures are given in Figure 5D.
Table 1. Comparison of the Crystal Structures, Optical Properties, and Efficiencies of Photocatalytic Degradation of MB Dye of the Zn$_2$SnO$_4$ Nanomaterials at Ambient Pressure Conditions and Their Recovered Counterparts after the Pressure Treatments

|                         | 0-D nanoparticles | 1-D nanostuctures |
|-------------------------|-------------------|-------------------|
| phase(s)                | I (inverse spinel)| I + HPOr (new high pressure orthorhombic) |
| absorption edge (nm); band gap/$E_g$ (eV) | ~353; 3.51 | ~322; 3.85 |
| main emission maxima (nm) | 358          | 328              |
| photodegradation of MB ($C/C_0$ in the presence of the samples after 35 min) | 0.3            | 0.78             |

$E_g$ orthorhombic: ∼3.37 eV) (Figure S3). The efficiency is, thus, found to vary according to the suitable band gap energy of the samples with respect to the light source ($E_g$ ≈ 3.39 eV). Therefore, a minor downturn in the degradation efficiencies is observed for the 0-D Zn$_2$SnO$_4$ nanoparticles after the pressure is released. In contrast, the degradation efficiencies are significantly enhanced for the recovered Zn$_2$SnO$_4$ nanostructures, where the heterojunction of the two phases, namely, inverse spinel ($E_g$: ∼4.19 eV) and orthorhombic phase ($E_g$: ∼3.37 eV), can effectively reduce the possibility of recombination of the photogenerated electrons and holes. The optical properties and efficiencies of photocatalytic degradation of the MB dye of the Zn$_2$SnO$_4$ nanomaterials at ambient pressure conditions and the recovered counterparts after the pressure treatments are assembled together in Table 1 mentioning their respective crystal structure.

## CONCLUSIONS

The impacts of the size and morphology of nanostructured Zn$_2$SnO$_4$ on their phase transformation behavior were investigated using 0-D Zn$_2$SnO$_4$ nanoparticles with an average particle size of ∼7 nm and 1-D Zn$_2$SnO$_4$ nanostructures with a typical chain-like morphology with several uneven interparticle voids. Unlike previously established martensitic phase transition of spinel structures, the 0-D Zn$_2$SnO$_4$ nanoparticles with an inverse spinel structure showed no significant pressure-induced structural modification under high pressure up to ∼40 GPa. On the other hand, an irreversible phase transformation was observed for the 1-D Zn$_2$SnO$_4$ inverse spinel Zn$_2$SnO$_4$ nanostructures to form a new quenchable high-pressure orthorhombic CaFe$_2$O$_4$-type structure. The inhomogeneously distributed pressure-induced strain relating to its distinct morphology is accounted for. Modifications of the optical band gaps and in emission properties can be seen after the pressure treatments. The photocatalytic degradation efficiencies of the original nanoparticles were compared to those of their respective pressure-recovered counterparts using the organic pollutant, MB. The pressure-recovered Zn$_2$SnO$_4$ nanostructures that have a heterojunction between the inverse spinel and the new orthorhombic phases appeared to be the most efficient photocatalyst.

## EXPERIMENTAL SECTION

The 0-D Zn$_2$SnO$_4$ nanoparticles of ∼7 nm size were prepared by the hydrothermal method, the most widely used technique for synthesizing nanomaterials. In contrast, the dense and elongated 1-D Zn$_2$SnO$_4$ nanostructures were prepared by calcining a stoichiometric mixture of the two presynthesized reactive precursors, namely, ZnO rod-like microstructures and SnO$_2$ nanoparticles. The explicit synthetic techniques for these precursors are mentioned elsewhere. The grain sizes and morphologies of the Zn$_2$SnO$_4$ nanostructures were examined by TEM analysis on a Tecnai G2 30ST. The HAADF-STEM images of the samples were taken using a JEOL JEM 2100F equipped with a CEG-corrector. The specific surface areas of the samples were measured using the BET method on a Quantachrome Instrument (iQ3, USA) by nitrogen adsorption–desorption studies at 77 K, whereas the BJJ method was followed for calculating the PSDs. XPS measurements were carried out in a PHI 5000 VersaProbe II scanning XPS microprobe (ULVAC-PHI, U.S.). High-pressure experiments were performed using a symmetric diamond anvil cell furnished with a pair of type-I diamonds with culet sizes of 300 μm. A preindentation stainless-steel gasket drilled with a 120 μm diameter hole at the center was used as the sample chamber, whereas Si oil was used as the pressure transmitting medium. A small ruby chip was used as the pressure gauge, and the R1-line emission of the ruby chip was used for pressure calibration during the measurements. In situ high pressure XRD measurements were conducted on the 6D beamline at Pohang Accelerator Laboratory (PAL) using a monochromatic 18.785 keV X-ray beam (λ = 0.6600 Å) with a diameter of 100 μm. A charge-coupled device detector was used for recording the Debye rings, which were further integrated into 1-D profiles using the Fit2D program. In situ high-pressure Raman measurements were performed by an inVia Raman spectrometer with a 532 nm excitation laser and 2400 g/cm grating at the Center for High Pressure Science & Technology Advanced Research, Shanghai, China, keeping the sample conditions similar to those used for the XRD measurements. The same spectrometer was used to measure the emission spectra of the samples using a UV laser of wavelength 325 nm (IK3301R-G He–Cd Laser, K KIMMON) as the excitation source. DR spectra of the samples were measured on a UV–vis–NIR spectrometer (Shimadzu UV-3600). For a typical experiment, the photocatalytic activities of the ambient and released samples were evaluated by measuring the degradation rate of MB (Merck, Germany) dye solution. An 8 W UV lamp (UVLS-28 EL Series) with an average light intensity of 0.6 mW/cm$^2$ and an emitting wavelength of 365 nm (UVC, ∼3.39 eV) was used as the irradiation source in a closed chamber. The dye degradation under UV light was periodically monitored by UV–vis spectroscopy (Shimadzu UV-3600).

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.9b01361.
Whole profile fitting of the XRD patterns of the 1-D Zn$_2$SnO$_4$ nanostructures at ~14, ~30, ~40 GPa, and after the pressure is released, scaled strain derived from the S–H equation based on the (111), (511), (220), and (440) diffraction planes of the 1-D Zn$_2$SnO$_4$ nanostructures, and absorption spectra of MB in the presence of the as-prepared and recovered 0-D Zn$_2$SnO$_4$ nanoparticles and 1-D Zn$_2$SnO$_4$ nanostructures under UV irradiation for different time intervals (PDF).

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**AUTHOR INFORMATION**

**Corresponding Author**
*E-mail: yongjaeelee@yonsei.ac.kr.*

**ORCID**

Partha Pratim Das: 0000-0001-6687-8324
P. Sujatha Devi: 0000-0002-6224-7821
Yongjae Lee: 0000-0002-2043-0804

**Notes**

The authors declare no competing financial interest.

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