

AFOSR Project Report

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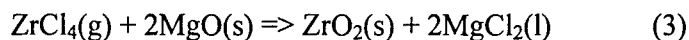
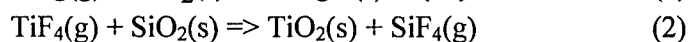
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14. ABSTRACT This project has been aimed at: 1) developing a better understanding of the manner in which the morphology and nanostructure of biologically-derived silica microtemplates evolve during the course of reactive conversion, and 2) determining which reaction parameters have the greatest impact on changes in morphology during such reactive conversion. The most significant accomplishments have been: - Demonstration (via high resolution TEM analyses) that the reaction of SiO₂ diatom frustules with Mg(g) at 650°C results in direct conversion into nanocrystalline MgO and Si (≤15 nm) without the formation of intermediate silicate phases. - Syntheses of MgO-converted frustules with minimal secondary phases (Si or Mg₂Si) by development of an optimized thermal treatment and selective dissolution treatment. - Identification and control of critical processing parameters to avoid active vaporization and gas-phase-assisted coarsening during the metathetic conversion of SiO₂ frustules into TiO₂. - Successful syntheses of ZrO₂ frustule replicas via a new two-step reaction process (conversion of SiO₂ into MgO via Mg(g) reaction; then conversion of MgO into ZrO₂ via ZrCl₄(g) metathetic reaction and selective MgCl₂ dissolution in water).					
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Objectives: 1) To develop a better understanding of the manner in which the morphology and nanostructure of biologically-derived silica microtemplates evolve during the course of reactive conversion, and 2) to determine which reaction parameters have the greatest impact on changes in morphology during such reactive conversion.

Summary of Effort: Work over the past year has focused on the conversion of silica-based diatom frustules by the net reactions:



where {Si} refers to Si in a Mg-Si liquid. Conversion of SiO₂-based diatom frustules into MgO, TiO₂, and ZrO₂ has been achieved with a preservation of the starting biogenic shapes.

Complete conversion of diatom frustules into MgO via net reaction (1) has been achieved at temperatures as low as 650°C within 2 h. Under these conditions, polycrystalline MgO frustules were generated with crystallite sizes of only 10-15 nm (i.e., much finer than for conversion at 900°C). TEM analyses of partially-converted specimens revealed the presence of MgO and Si nanocrystals (<5 nm in dia.) in close proximity within an unreacted silica matrix; that is, the conversion of SiO₂ appears to proceed directly without the formation of intermediate silicate phases (as has been reported in the literature for the reaction of Mg(g) gas with bulk SiO₂ at 1150°C). The solid Si product then reacted with excess Mg(g) to form Mg₂Si. The amount of Mg₂Si generated by this latter reaction was minimized by controlling the Mg:SiO₂ reactant ratio and the reaction temperature and time. A sodium hydroxide solution was then used to selectively dissolve away the Si product of reaction (1) to yield MgO-based frustule replicas.

Complete conversion of SiO₂-based diatom frustules into TiO₂ via the net reaction (2) has been accomplished at only 350°C via a two-step process: i) the formation of TiOF₂(s) via reaction with TiF₄(g) and then ii) conversion of TiOF₂(s) into TiO₂(s) via reaction with O₂(g). By controlling the TiF₄:SiO₂ reactant ratio, the heating rate, and the peak reaction temperature and time during the first stage of reaction, active vaporization (due to TiOF₂(s) formation away from the frustule surface) and coarsening (via a gas phase diffusion mechanism) were minimized so as to yield TiOF₂ replicas that preserved the starting frustule shape and fine features.

A two-step reaction process has also been developed to allow for chemical conversion of the silica-based diatom frustules into zirconia. Initial attempts to form ZrO₂ replicas via direct reaction of ZrF₄(g) with SiO₂(s) resulted in disintegration of the frustules. To avoid the formation of volatile Si-O-bearing products associated with such disintegration, the SiO₂ frustules were first converted into MgO via net reaction (1). The MgO replicas were then exposed to ZrCl₄(g) at 650°C for 2 h to allow for reaction (3). The latter reaction yielded tetragonal ZrO₂ replicas along with a liquid MgCl₂ product that then solidified during cooling. Selective dissolution of the solidified magnesium chloride in water then yielded free-standing zirconia frustule replicas.

The reaction-based processes developed in this BioInspired Concepts (BIC) project have enabled the syntheses of frustule-shaped structures that exhibit:

- i) luminescence (from Eu/BaTiO₃ coatings applied to compatible MgO-converted frustules)
- ii) gas sensing (due to rapid changes in the conductivity of TiO₂ frustules resulting from small changes in ethanol vapor pressure)
- iii) organophosphor destruction (pesticide-bearing solutions have exhibited rapid reaction with TiO₂-converted frustules)

Accomplishments/New Findings:

- Demonstration (via high resolution TEM analyses) that the reaction of SiO₂ diatom frustules with Mg(g) at 650°C results in direct conversion into nanocrystalline MgO and Si (≤15 nm) without the formation of intermediate silicate phases.
- Syntheses of MgO-converted frustules with minimal secondary phases (Si or Mg₂Si) by development of an optimized thermal treatment and selective dissolution treatment.
- Identification and control of critical processing parameters to avoid active vaporization and gas-phase-assisted coarsening during the metathetic conversion of SiO₂ frustules into TiO₂.
- Successful syntheses of ZrO₂ frustule replicas via a new two-step reaction process (conversion of SiO₂ into MgO via Mg(g) reaction; then conversion of MgO into ZrO₂ via ZrCl₄(g) metathetic reaction and selective MgCl₂ dissolution in water).

These new accomplishments have made possible the development (in a MURI project) of luminescent diatom frustules (for microtagging applications), single diatom frustule gas sensors (for minimally-invasive sensing), and diatom frustules that rapidly destroy organophosphorous compounds (potentially for removing chemical warfare agents).

Reactive Conversion into MgO Frustules

Phase and Nanostructural Evolution at 650°C

In order to slow down the rate of reaction (to enable evaluation of the earliest stage(s) of the MgO conversion process) *Aulacoseira* diatom frustules were exposed to Mg(g) at only 650-700°C. The frustules were placed inside and near one end of a mild steel tube. Mg granules (99.8% purity, 0.3-1.7 mm size) were placed at the other end of the tube. An excess of Mg granules relative to silica frustules (Mg:SiO₂ molar ratio = 9.9:1) was placed inside the steel tubes in order to ensure that the source of magnesium vapor was not depleted during the course of the reaction. The tube was then crimped and welded shut within an Ar atmosphere inside a glove box. The resulting sealed steel ampoule was then bent into an inverted "V" shape, in order to ensure that the silica frustules remained physically separated from the magnesium powder. The sealed steel ampoules were then heated at 5-7°C/min to 650°C or 700°C and held for 0.5-2 h.

A secondary electron image of a starting *Aulacoseira* diatom frustule, and an associated energy-dispersive x-ray (EDX) pattern are shown in Figures 1a and 1b, respectively (note: the gold peak was obtained from a thin coating applied to avoid electron beam charging).

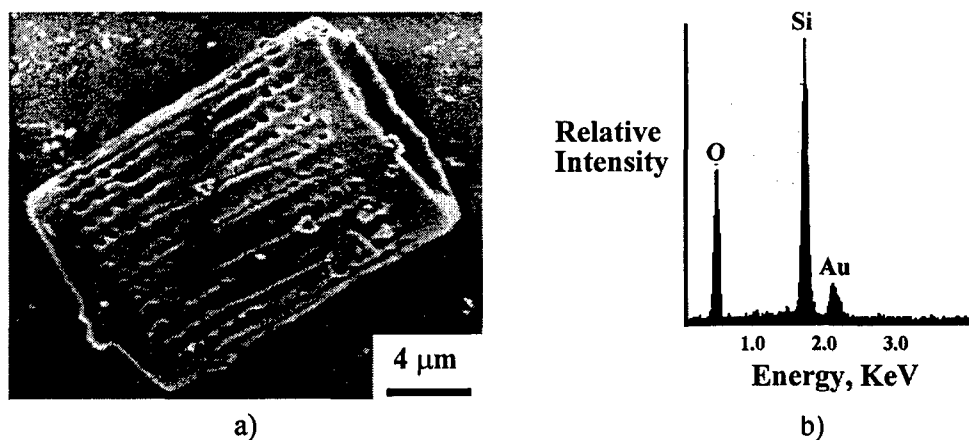


Fig. 1. a) Secondary electron image and b) EDX analysis of an *Aulacoseira* diatom frustule.

Upon heating to 650-700°C, the silica frustules within the sealed steel tube were exposed to magnesium gas generated by evaporation from the magnesium granules. Magnesium gas may undergo the following oxidation-reduction (displacement) reaction with silica:



A secondary electron image of an *Aulacoseira* frustule that had been exposed to Mg(g) for 0.5 h at 700°C is shown in Fig. 2a. The cylindrical shape, narrow channels, and rows of fine pores were well preserved in this reacted frustule. A TEM image of a cross-section of such a reacted frustule is shown in Fig. 2b. The cross-section was comprised largely of crystallites with diameters ≤ 50 nm, although a few larger crystallites (100-150 nm diameter) were also detected. EDX (Fig. 2c) and electron diffraction (Fig. 2d) analyses revealed that the reacted frustule was comprised of magnesium oxide (periclase), with very little silicon detected throughout the cross-section (the gold peak in Fig. 2c was obtained from a coating applied to the specimen for SEM analysis prior to ion milling; the copper peak was obtained from the grid used to support the specimen cross-section in the TEM). The loss of silicon was a result of the formation of a Mg-Si

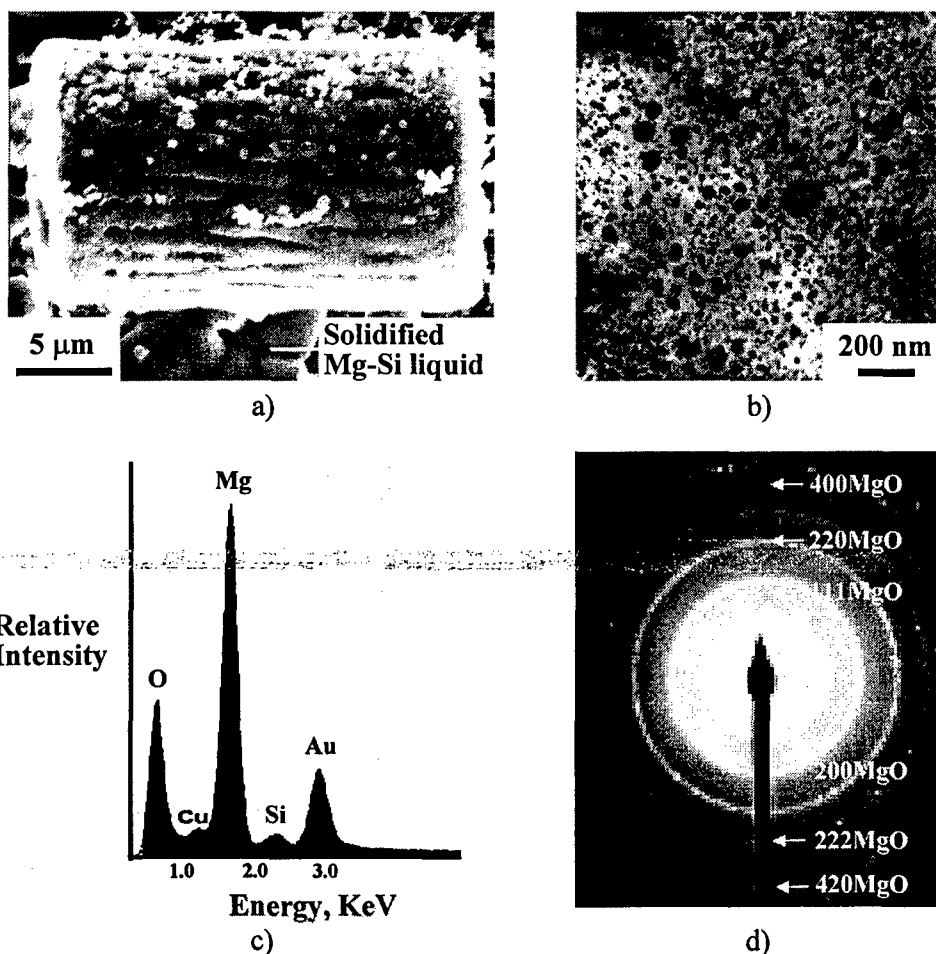


Fig. 2. a) Secondary electron image of an *Aulacoseira* frustule after exposure to Mg(g) for 0.5 h at 700°C. b) Transmission electron microscope image, c) EDX analysis, and d) electron diffraction analysis of an electron transparent cross-section of an *Aulacoseira* frustule after exposure to Mg(g) for 0.5 h at 700°C.

liquid (a eutectic liquid can form at 638°C in the Mg-Si system) that poured out of the frustule. The solidified Mg-Si liquid can be seen below the MgO-converted frustule in Fig. 2a. The data in Fig. 2 indicate that *Aulacoseira* frustules can be fully converted into a nanocrystalline MgO-based replica within only 0.5 h of exposure to Mg(g) at 700°C.

In order to evaluate the phase and nanostructural evolution during such shape-preserving reactive conversion, the reaction temperature was further reduced to 650°C to slow the rate of reaction. XRD analyses of the *Aulacoseira* diatom frustules before and after exposure to Mg(g) for various times at 650°C are shown in Fig. 3. For exposure times of 1 h or less, only diffraction peaks for SiO₂ (as cristobalite) were detected. However, from 1.5 to 2 h of exposure to Mg(g), MgO, Si and Mg₂Si formed at the expense of cristobalite. Indeed, after the 2 h treatment, the diffraction peaks for cristobalite vanished. Bright field TEM images, and associated electron diffraction patterns, of ion-milled cross-sections obtained from specimens exposed to Mg(g) for 1-2 h at 650°C are shown in Figs. 4a-f. Consistent with XRD analyses, selected area electron diffraction (SAED) analyses of the cross-section of the 1 h specimen in Fig. 4a revealed only diffraction associated with cristobalite (hkl values associated with

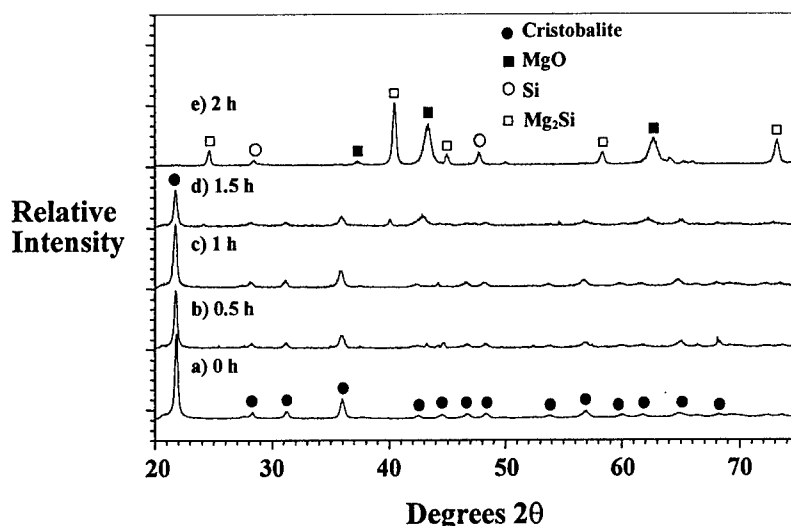


Fig. 3. XRD analyses of *Aulacoseira* diatom frustules a) before and b)-e) after exposure to magnesium gas at 650°C for 0.5, 1, 1.5 and 2 h.

several of these diffraction spots are indicated). Prolonged exposure of this specimen cross-section to the electron beam in the TEM also resulted in amorphization, as has been observed by other authors during TEM analyses of silica-based specimens. After 1.5 h of reaction, the specimen cross-section was found to be partially reacted (Fig. 4b). SAED analysis of the reaction zone (Fig. 4e) revealed the presence of MgO and Si, consistent with reaction (1). The absence of electron or x-ray diffraction peaks (Figs. 3d and 4e) for magnesium silicates (e.g., forsterite, Mg₂SiO₄; enstatite, MgSiO₃) suggested that these compounds were not involved as intermediate products in reaction (1). Consistent with such diffraction analyses, a high-resolution TEM image (Fig. 5) of this cross-section obtained at the location of the interface between the reacted zone and the unreacted silica revealed the presence of only MgO and Si nanocrystals (≤ 5 nm in diameter) in an amorphous matrix. The structure generated after 2 h of exposure to Mg(g) at 650°C is shown in Fig. 4c. The nanocrystalline reaction zone completely penetrated this specimen. SAED analysis revealed a mixture of MgO, Si, and Mg₂Si in the

reaction zone. The latter compound was generated by the continued reaction of the silicon product from reaction (1) with magnesium gas, as shown below.



As discussed above, further reaction of this Mg_2Si product with Mg(g) leads to the formation of a Mg-Si liquid that tends to pour out of the converted frustule onto the underlying steel substrate (Fig. 2a), so as to leave a nanocrystalline MgO-based structure. The width of the most intense x-ray diffraction peak for magnesia (the (200) peak near $2\theta = 43$ degrees) was obtained from several batches of specimens exposed to the $650^\circ\text{C}/2$ h heat treatment. Insertion of this data into the Scherrer equation yielded average magnesia crystallite sizes of 12-15 nm. These XRD-derived values were consistent with the fine nanocrystallites observed in the TEM images of Figs. 4b and c. These MgO grains were also considerably finer, by several orders of magnitude, than those generated upon exposure of diatom silica to magnesium gas at 900°C for 4 h.

Prior work by other authors on the reaction of sintered $\text{SiO}_2(\text{s})$ tablets or quartz crystals with Mg(g) at much higher temperatures (e.g., 1150°C) have indicated that $\text{Mg}_2\text{SiO}_4(\text{s})$ and SiO(g)

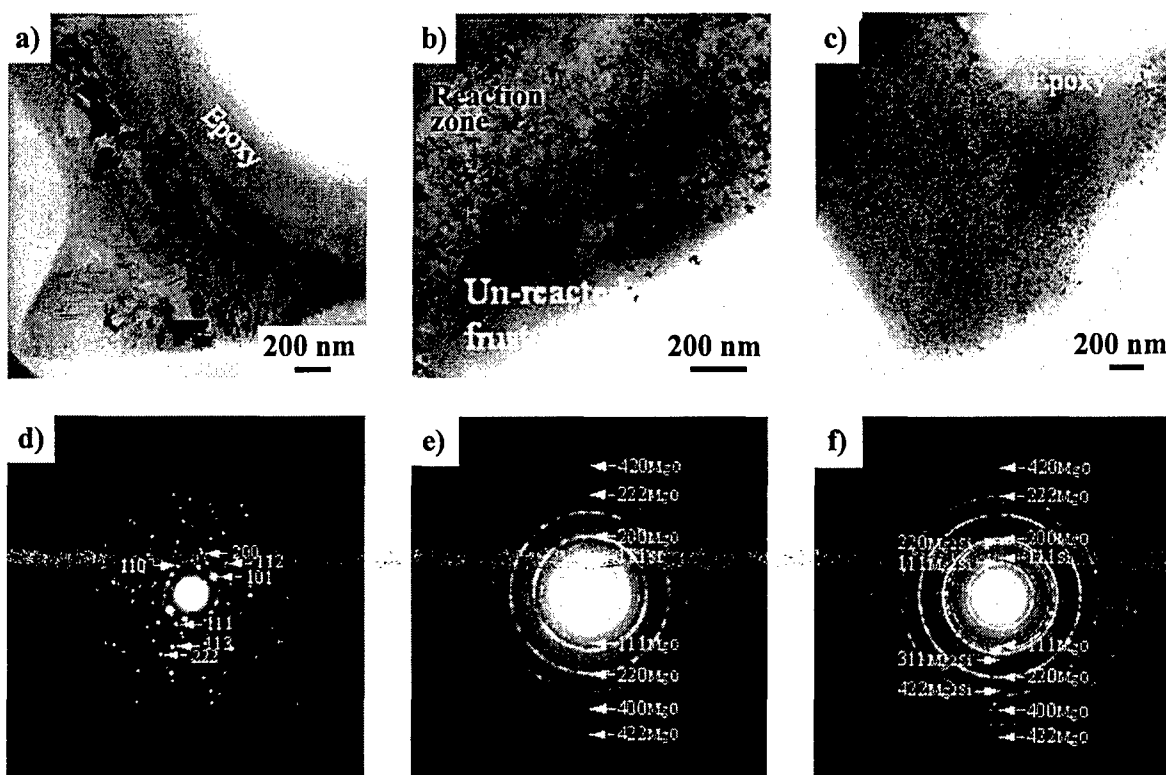
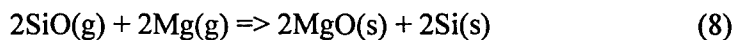


Fig. 4. a)-c) TEM images of cross-sections of *Aulacoseira* frustules exposed to Mg gas at 650°C for 1, 1.5, and 2 h respectively. d)-f) SAED patterns obtained from a)-c) respectively.

formed as intermediate reaction products via the following reaction steps:



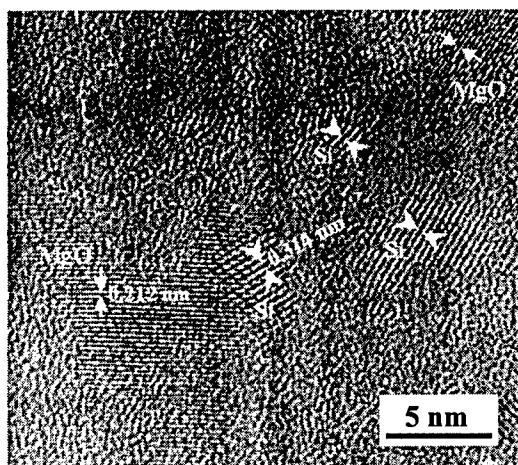


Fig. 5. High-resolution TEM image of a cross-section of an *Aulacoseira* diatom frustule exposed to Mg gas at 650°C for 1.5 h (obtained at the reaction front seen in Fig. 4b).

While the formation of $\text{Mg}_2\text{SiO}_4(\text{s})$ and $\text{SiO}(\text{g})$ via reaction (6) was thermodynamically favored at 1150°C, this reaction was not favored at 650°C in the present work. The thermodynamic ratio of the $\text{Mg}(\text{g})$ to $\text{SiO}(\text{g})$ partial pressures that is required for reaction (6) to proceed spontaneously to the right at 650°C is 48, compared to only 6.5×10^{-2} at 1150°C. The amount of Mg relative to SiO_2 that was sealed in the steel ampoules in the present work (a molar $\text{Mg}:\text{SiO}_2$ ratio of 9.9:1) was insufficient to allow for the occurrence of this intermediate reaction.

The basic understanding of the phase and nanostructural evolution obtained from the work discussed above has enabled the development of optimized processing treatments over the past year for the conversion of SiO_2 frustules into MgO-based frustules containing minimal secondary phases (i.e., Si or Mg_2Si). By minimizing these secondary phases, the MgO frustules could be utilized as chemically-compatible substrates for other functional ceramic coatings (e.g., luminescent Eu-doped BaTiO_3) and as structures for microfluidic mixing devices (e.g., based on electroosmotic flow). Optimization of the MgO conversion conditions is discussed below.

Optimized Processing of MgO-converted Frustules

Several processing changes have been introduced in order to reduce the relative amounts of undesired second phases (e.g., Si, Mg_2Si) in the MgO-converted frustules. A schematic of a typical reaction configuration within the sealed iron tubes is shown in Fig. 6a. At the start of the experiments, the diatom frustules and solid magnesium reactants were placed at opposite ends of the iron tube. The tube was then crimped in the middle (to avoid physical mixing of the reactants) and the ends were welded shut. Upon heating the magnesium vaporized and diffused to the frustules, whereupon the displacement reaction commenced. As shown in Figs. 6a and 6b, distinct reaction zones were detected within partially-converted frustule powder beds. The distinct interfaces and colors of these zones enabled separation and analyses (XRD, SEM, EDX) of the phases within each zone. The region located nearest the magnesium gas source was observed to possess a grey or blue-grey color, which was a result of the presence of Mg_2Si . A secondary electron image of this phase is shown in Fig. 6c. The darker intermediate zone was comprised of a mixture of MgO and Si. A secondary electron image of a frustule from within this zone, exhibiting Si nodules, is shown in Fig. 6d. The brown zone at the opposite end of the

powder bed contained residual unreacted silica. The distribution of these products was consistent with the reaction steps identified above; that is, SiO_2 is first converted into a mixture of MgO and Si , and the Si then reacts with excess $\text{Mg}(\text{g})$ to form Mg_2Si .

The $\text{Mg}:\text{SiO}_2$ reactant ratio, the reaction temperature, and the reaction time were varied to allow for complete conversion of the SiO_2 into MgO while minimizing the amount of Mg_2Si produced. With a $\text{Mg}:\text{SiO}_2$ ratio of 2:1 and a reaction temperature/time of $900^\circ\text{C}/1.5$ h, the frustule powder bed was converted into a mixture of MgO and Si with minimal Mg_2Si , as shown in Fig. 7a. The residual silicon was then selectively dissolved away with an aqueous sodium hydroxide solution. The NaOH in this solution was adjusted to achieve a $\text{NaOH}:\text{Si}$ molar ratio of 2:1 to allow for the formation of soluble sodium metasilicate (Na_2SiO_3). The Si -depleted frustules were then rinsed with boiling DI water. XRD analysis and secondary electron images of the resulting MgO frustules are shown in Figs. 7b and 8, respectively. It is important to note that the development of processing methods for removing Mg_2Si (through adjustment of reaction conditions) and Si (through selective dissolution) were essential for the use of MgO frustules as: i) substrates for conformal luminescent coatings (i.e., Eu -doped BaTiO_3 coatings on MgO), and ii) 3-D micro-structures for incorporation in electro-osmotic mixing devices (i.e., to avoid bubble formation due to the reaction of residual Mg_2Si with water).

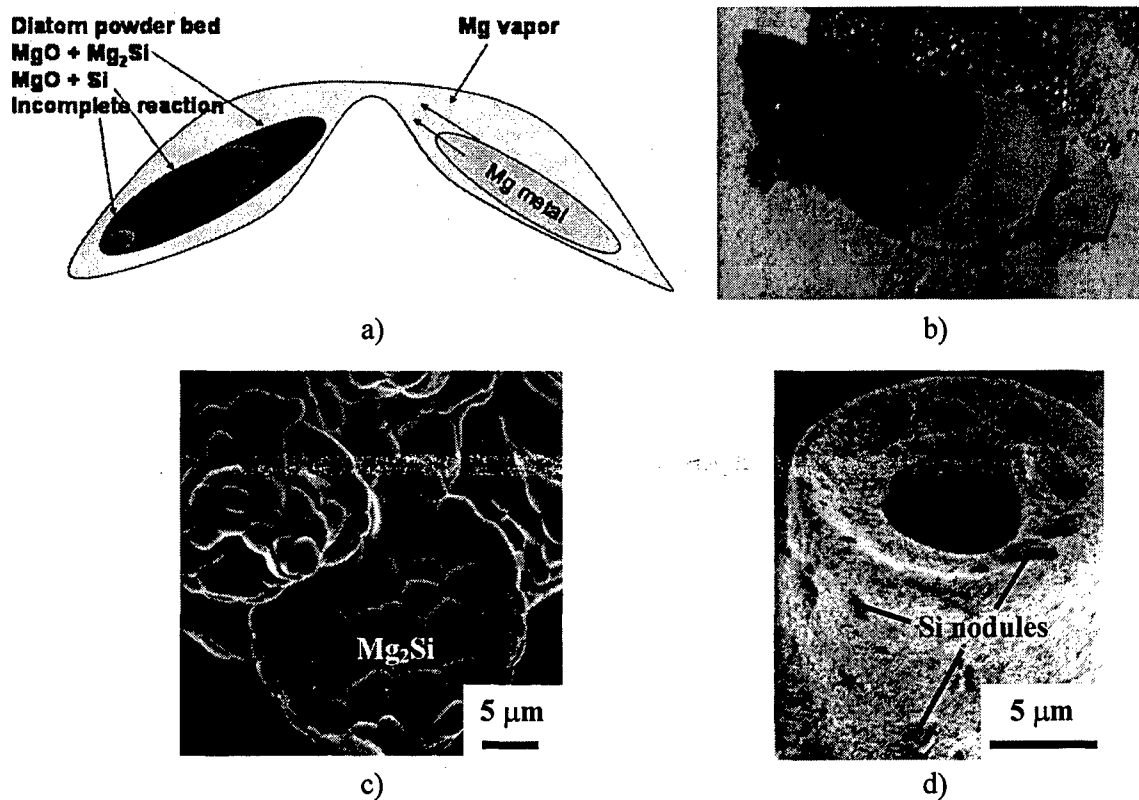


Fig. 6. a) Schematic illustration of the sealed iron reaction tube showing the positioning of reactants and the typical progression of products through the frustule powder bed. b) Optical image of a frustule powder bed after reaction for 3 h at 850°C . Secondary electron images revealing the Mg_2Si and Si secondary phases are shown in c) and d), respectively.

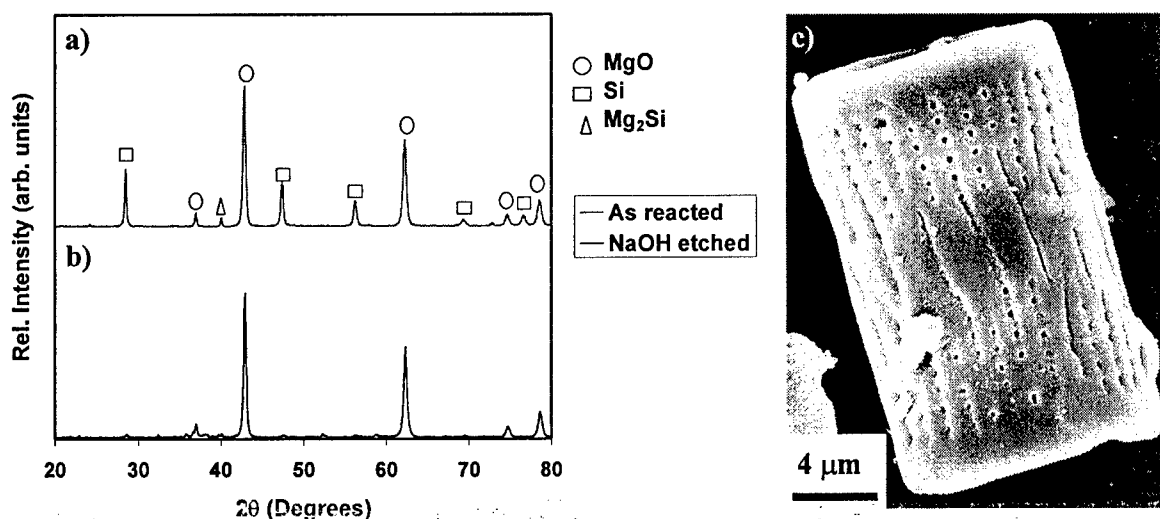


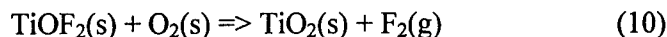
Fig. 7. XRD analyses of MgO-converted diatoms a) before and b) after selective silicon dissolution with an aqueous sodium hydroxide solution. c) Secondary electron image of the MgO frustule corresponding to the XRD pattern in b).

Reactive Conversion into TiO₂ (anatase) Frustules

A two-step reaction process has been used to convert silica-based diatom frustules into titania replicas. The first step involves the following metathetic reaction of SiO₂ with TiF₄(g) to yield the oxyfluoride, TiOF₂:



In the second step, the TiOF₂ is converted to TiO₂ by reaction with O₂(g):



In this work, solid TiF₄ was utilized as a low-temperature source of TiF₄ vapor (note: the sublimation temperature of TiF₄(s) is 285°C). 100 milligrams of *Aulacoseira* diatom frustules were placed within a titanium tube along with solid TiF₄. As discussed above for the Mg/SiO₂ reactions, both ends of the tube were then crimped and welded shut. The specimen-bearing tube was then heated to the desired reaction temperature. At the end of this treatment, the tube was pushed out of the hot zone of the horizontal tube furnace. For the second reaction, the specimens were heated to 350°C for 2 h in pure, flowing oxygen.

Several processing parameters associated with the first reaction step have been found to be particularly critical with regards to shape preservation:

- i) the TiF₄:SiO₂ reactant ratio
- ii) the peak reaction temperature and time
- iii) the heating rate

Initial experiments conducted at 500-700°C for 2 h using TiF₄:SiO₂ molar ratios of $\geq 4.9:1$ within the sealed Ti tubes resulted in disintegration of the starting SiO₂ frustules and the formation of relatively coarse plate-shaped TiO₂ crystals (5-10 μm X 5-10 μm X 0.5-1 μm thick). Such active vaporization of the silica frustules was avoided by lowering the reaction temperature to 350°C and reducing the TiF₄:SiO₂ molar ratio to 2.4:1 within the Ti tubes. After 2 h under these conditions, the overall shape and fine features of the reacted frustule were quite similar to those

of the starting frustules, as shown in Fig. 8a below. EDX and XRD analyses confirmed that the frustules had been converted completely into TiOF_2 . Further reduction in the $\text{TiF}_4\text{:SiO}_2$ molar ratio to 1.6 resulted in negligible reaction of the frustules within 2 h at 350°C .

An increase in the reaction temperature from 350°C to 450°C for a fixed time (2 h), a fixed heating rate ($5^\circ\text{C}/\text{min}$), and a fixed $\text{TiF}_4\text{:SiO}_2$ ratio resulted in appreciable coarsening of the TiOF_2 grains, as shown in Figs. 8a and 8b. For a constant peak temperature/time of $350^\circ\text{C}/2$ h, a reduction in the heating rate also yielded an increase in TiOF_2 crystallite size (Figs. 8c and 8d). At such modest temperatures and times, it is not likely that such coarsening occurred by a solid-state diffusion mechanism. Given that a gaseous TiOF_2 species can exist at these temperatures, it is likely that such rapid coarsening occurred via transport through the gas phase; that is, by the evaporation of small TiOF_2 particles and condensation onto larger TiOF_2 particles (Ostwald ripening) as illustrated in Fig. 8e. Such gas-phase-assisted coarsening has been observed in a number of ceramic systems with volatile species (notably by D. W. Readey: "Vapor Transport and Microstructure Development in Ceramics," *Journal of Metals*, 36 (12) 18-18, 1984).

By controlling the heating rate, peak reaction temperature and time, and $\text{TiF}_4\text{:SiO}_2$ molar ratio, the active vaporization of silica and the rapid vapor-phase-assisted coarsening of TiOF_2 could be minimized so as to yield TiOF_2 and then TiO_2 replicas that retained the shapes and fine features of the starting diatom frustules.

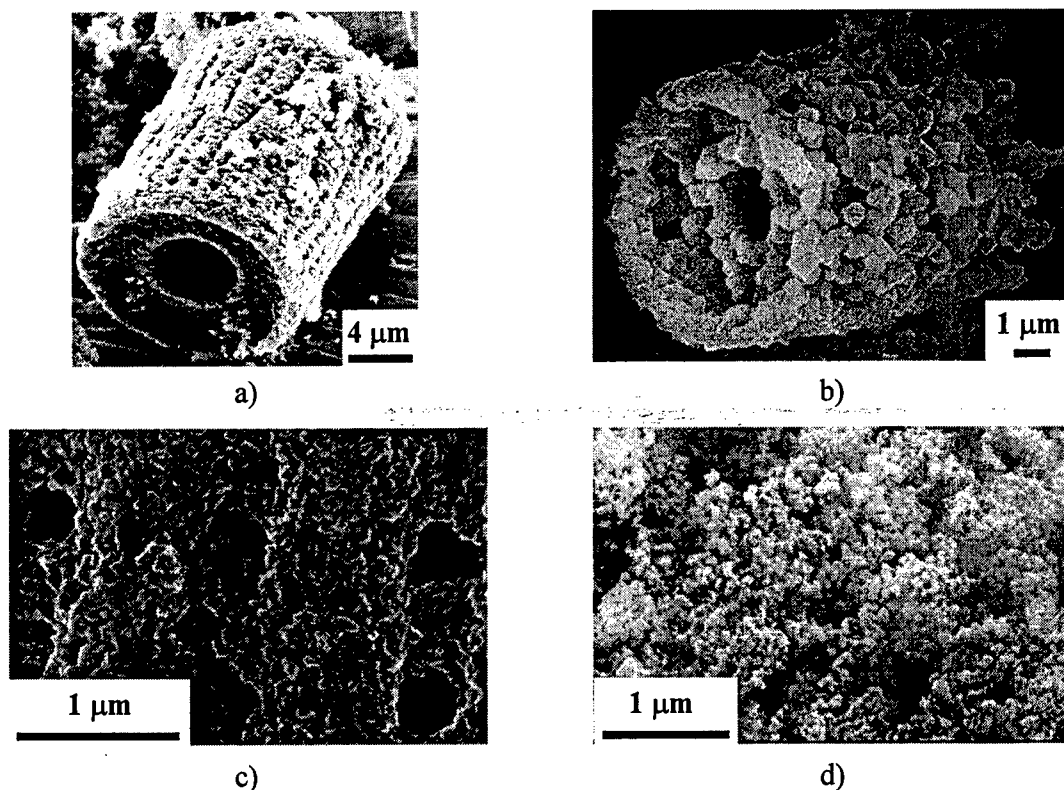


Fig. 8. Secondary electron images of TiOF_2 -converted frustules generated by reaction for 2 h at: a) 350°C (heating rate = $5^\circ\text{C}/\text{min}$), b) 450°C (heating rate = $5^\circ\text{C}/\text{min}$). Higher magnification images of TiOF_2 -converted frustules after reaction for 2 h at 350°C with heating: c) at a rate of $5^\circ\text{C}/\text{min}$ and d) via thrusting into pre-heated furnace.

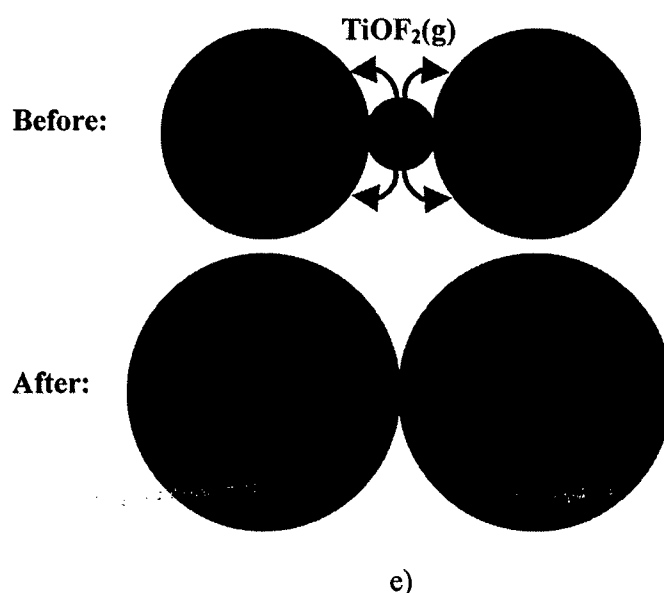


Fig. 8 (cont.). A schematic illustrating gas-phase-assisted coarsening is shown in e).

Reactive Conversion into ZrO_2 Frustules

The direct reaction of $\text{ZrF}_4(\text{g})$ with SiO_2 diatom frustules was conducted within sealed nickel alloy tubes for 2 h over a range of temperatures (250-800°C) and $\text{ZrF}_4\text{:SiO}_2$ molar ratios (0.36 to 3.6). For reaction temperatures at or below 350°C, no reaction was detected within 2 h. At higher temperatures and for all $\text{ZrF}_4\text{:SiO}_2$ ratios examined, the silica frustules were observed to disintegrate, as shown in Fig. 9.

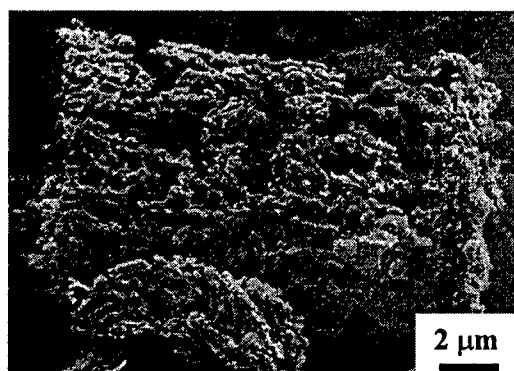
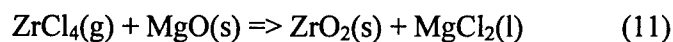


Fig. 9. Secondary electron image of a disintegrating *Aulacoseira* frustule after exposure to ZrF_4 gas at 600°C for 2 h.

An entirely different two-step reaction process was then adopted so as to avoid such active vaporization. In the first step, the silica frustules were converted into MgO , as per the oxidation-reduction conditions discussed above. The magnesia was then exposed to $\text{ZrCl}_4(\text{g})$ to allow for the following net reaction:



The magnesium chloride product, which solidified upon cooling to room temperature, was then be removed by selective dissolution in water. To test this new approach, the magnesia-converted frustules were sealed along with zirconium chloride ($\text{ZrCl}_4\text{:MgO}$ molar ratio = 5.8) in nickel alloy tubes. The tubes were then heated at $5^\circ\text{C}/\text{min}$ to a peak temperature of $325\text{--}650^\circ\text{C}$ for 2 h. The reacted specimens were then immersed and stirred in water heated to 80°C for 1 h. For reaction temperatures at or below 475°C , significant amounts of residual unreacted magnesium oxide were detected. However, reaction for 2 h at 650°C resulted in complete conversion of the magnesia into tetragonal zirconia and magnesium chloride. An XRD pattern of such reacted specimens (after dissolution of the magnesium chloride in water) is shown in Fig. 10. Secondary electron images of zirconia-converted frustules and a corresponding EDX pattern are shown in Figs. 11a-c. The zirconia replica retained the cylindrical shape of the starting *Aulacoseira* frustule.

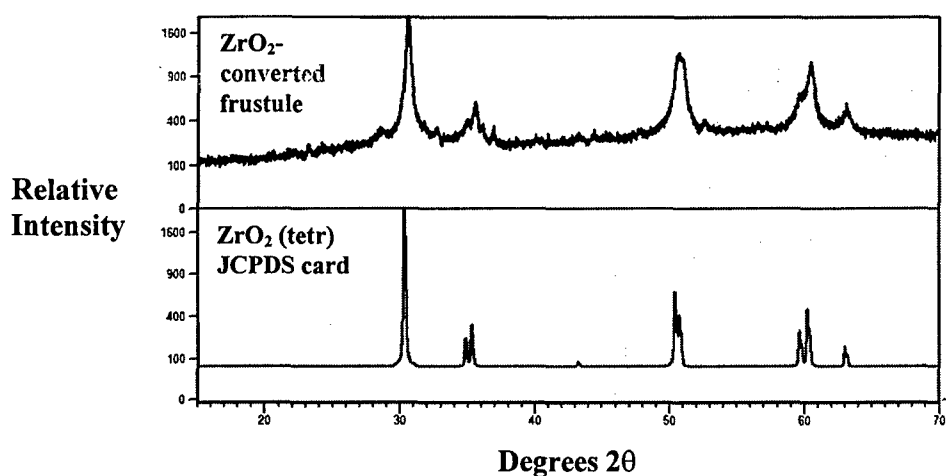
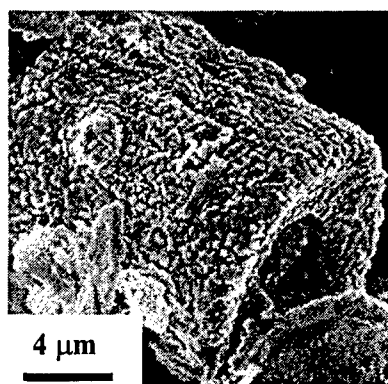
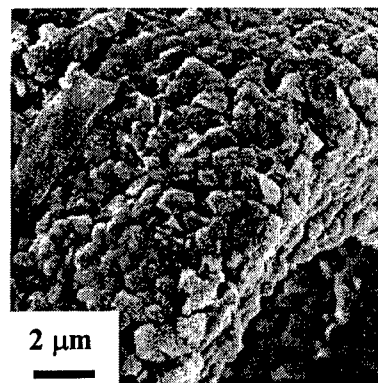


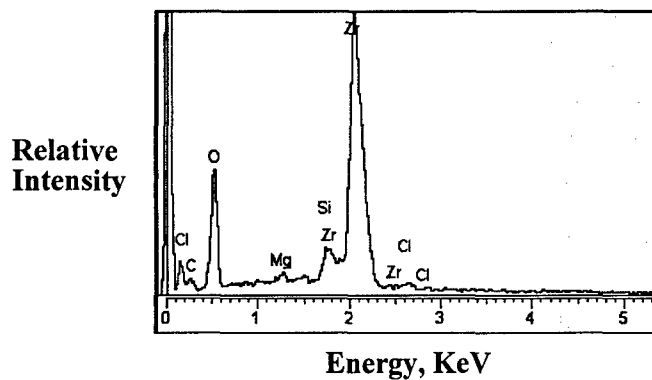
Fig. 10. XRD pattern obtained from a ZrO_2 -converted frustule generated by exposure of a MgO -converted frustule to $\text{ZrCl}_4(\text{g})$ for 2 h at 650°C followed by immersion in stirred water at 80°C for 1 h.



a)



b)



c)

Fig. 11. a), b) Secondary electron images, and c) an associated EDX pattern obtained from a ZrO_2 -converted frustule generated by exposure of a MgO -converted frustule to $\text{ZrCl}_4(\text{g})$ for 2 h at 650°C followed by immersion in stirred water at 80°C for 1 h.

Personnel Supported:

Faculty: Prof. Ken H. Sandhage
Graduate Students: Mr. Samuel Shian
Mr. Shawn Allan
Mr. Matthew Dickerson

Publications:

1. Y. Cai*, S. M. Allan, F. M. Zalar, K. H. Sandhage, "Three-dimensional Magnesia-based Nanocrystal Assemblies via Low-Temperature Magnesiothermic Reaction of Diatom Microshells," *J. Am. Ceram. Soc.*, **88** [7] 2005-2010 (2005).
2. M. B. Dickerson, R. R. Naik, P. M. Sarosi, G. Agarwal, M. O. Stone, K. H. Sandhage, "Ceramic Nanoparticle Assemblies with Tailored Shapes and Tailored Chemistries via Biosculpting and Shape-preserving Inorganic Conversion," *J. Nanosci. Nanotech.*, **5** [1] 63-67 (2005).
3. M. R. Weatherspoon*, S. M. Allan, E. Hunt*, Y. Cai*, K. H. Sandhage, "Sol-Gel Synthesis on Self-Replicating Single-Cell Scaffolds: Applying Complex Chemistries to Nature's 3-D Nanostructured Templates," *Chem. Comm.*, [5] 651-653 (2005).
4. M. B. Dickerson, R. R. Naik, M. O. Stone, Y. Cai*, and K. H. Sandhage, "Identification of Peptides that Promote the Rapid Precipitation of Germania Nanoparticle Networks via Use of a Peptide Display Library," *Chem. Comm.*, **15**, 1776-1777 (2004).
5. R. R. Unocic, F. M. Zalar, Peter M. Sarosi, Ye Cai*, and K. H. Sandhage, "Anatase Assemblies from Algae: Coupling Biological Self-assembly of 3-D Nanoparticle Structures with Synthetic Reaction Chemistry," *Chem. Comm.*, [7] 795-796 (2004).

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Interactions/Transitions:

The Sandhage group has moved from Ohio State University to the Georgia Institute of Technology. New facilities that have been set up at Georgia Tech for the Sandhage group over the past year include:

- a diatom culturing facility (with the advice and help of M. Hildebrand, UCSD/Scripps)
- a biopanning facility (with the advice and help of R. Naik, AFRL)
- new thermal processing facilities (with the advice and help of R. Rapp, Ohio State)

A new state-of-the-art x-ray characterization facility (which includes a high-temperature x-ray diffraction system for the work in this BIC program) has also been established over the past year by R. Snyder, a collaborator with K. Sandhage.

The reaction-based processes developed in this BioInspired Concepts (BIC) project have enabled the syntheses of frustule-shaped structures (via a MURI project) that exhibit:

- i) luminescence (from Eu/BaTiO₃ coatings applied to compatible MgO-converted frustules)
- ii) gas sensing (due to rapid changes in the conductivity of TiO₂ frustules resulting from small changes in ethanol vapor pressure)
- iii) organophosphor destruction (pesticide-bearing solutions have exhibited rapid reaction with TiO₂-converted frustules)

Luminescence measurements have been obtained via a new collaboration with Prof. Chris Summers of the MSE Dept. at Georgia Tech. Gas sensing measurements have been obtained via a new collaboration with Prof. Meilin Liu of the MSE Dept. at Georgia Tech. Organophosphor destruction kinetics have been measured in a new collaboration with Prof. Ching-Hua Huang of the Civil and Environmental Engineering Dept. at Georgia Tech.

Other interactions include the following technical presentations:

1. M. S. Haluska, K. H. Sandhage, R. L. Snyder, S. T. Misture, "High Temperature In-Situ X-ray Diffraction of the $\text{Mg(g)}/\text{SiO}_2\text{(s)}$ Diatom Displacement Reaction," 53rd Annual Denver X-ray Conference, Denver, CO, August 3, 2004.
2. R. L. Snyder, K. H. Sandhage, "A Novel Biology-based Route to Three-Dimensional Nanoparticle Assemblies with Intricate Shapes, Nanoscale Features, and Tailored Chemistries: Bioclastic and Shape-preserving Inorganic Conversion (BaSIC)," Plenary talk, 7th International Workshop on High-Temperature Superconductors and Novel Inorganic Materials Engineering, Moscow, Russia, June 20-25, 2004.
3. K. H. Sandhage, C. Gaddis, M. Dickerson, S. Allan, P. Graham, S. Shian, M. Weatherspoon, Y. Cai, R. R. Naik, M. O. Stone, "A Hybrid (Biogenic/Synthetic) Route to 3-D Nanoparticle Assemblies with Tailored Chemistries: the Bioclastic and Shape-preserving Inorganic Conversion (BaSIC) Process," Invited talk, MRS Spring Meeting, San Francisco, CA, March 12-16, 2004.
4. S. Allan, M. Dickerson, C. Gaddis, S. Shian, R. R. Unocic, F. M. Zalar, Y. Cai, E. Hunt, K. H. Sandhage, R. R. Naik, M. O. Stone, M. Hildebrand, B.P. Palenik, "3-D Nanoparticle Structures with Self-Replicating Shapes and Synthetically-Tailored Compositions," 133rd Annual TMS Meeting, Charlotte, NC, March 16, 2004.
5. K. H. Sandhage, C. S. Gaddis, M. B. Dickerson, S. Shian, R. R. Naik, M. O. Stone, M. M. Hildebrand, B. P. Palenik, "The BaSIC Route to 3-D Nanoparticle Assemblies with Self-Replicating, Genetically-Tailored Shapes and Synthetically-Tailored Chemistries," 28th International Cocoa Beach Conf. and Exposition on Advanced Ceramics and Composites, Cocoa Beach, FL, January 27, 2004.
6. C. S. Gaddis, J. Zhao, K. H. Sandhage, "A Transient Bioscaffolding Route to Chemically-Tailored, Nanoparticle-based Assemblies with Complex 3-D Shapes and Fine (Meso-to-Nanoscale) Features," Materials Research Society Fall Meeting, Boston, MA, December 2, 2003.
7. R. R. Unocic, F. M. Zalar, K. H. Sandhage, "A Hybrid (Biogenic/Synthetic) Route to 3-D Nanoparticle Assemblies with Tailored Chemistries: The Bioclastic and Shape-preserving Inorganic Conversion (BaSIC) Process," Materials Research Society Fall Meeting, Boston, MA, December 2, 2003.
8. K. H. Sandhage, M. B. Dickerson, P. M. Sarosi, G. Agarwal, R. R. Naik, M. O. Stone, "Chemical Transformation of Diatoms," Invited talk, Diatom Nanotechnology Workshop at the North American Diatom Society meeting, Florida Keys, October 22-26, 2003.
9. K. H. Sandhage, "Chemically-Tailored 3-D Nanoparticle Structures by the BaSIC (Bioclastic and Shape-preserving Inorganic Conversion) Process," Invited talk, Composites at Lake Louise Conference, Lake Louise, Canada, October 21, 2003.