Solution Synthesis of Spinel



Ceramic, Composite and Optical Materials Center Rutgers, The State University of New Jersey

Richard Riman, John Wilson, and Daniel Kopp

Background

- Spinel (MgAl₂O₄) is a very attractive material for the preparation of IR-transparent windows that have excellent ballistic protection properties.
- Current approaches for commercially producing spinel use high temperature (600 to 1000°C) solid-state reaction methods along with subsequent milling, leading to reduced purity and high energy costs.
- Only one manufacturer of high-quality spinel powder in the USA, Baikowski and the cost is high (\$60/kg).*
- There is an opportunity to research, develop and commercialize low-energy hydrothermal technology that can produce low-cost spinel.





Overall Goals

- Find the mildest reaction conditions and shortest reaction times to prepare MgAl₂O₄ spinel powder direct from solution (with <u>no</u> high temperature treatment).
- Study the crystallization and reaction kinetics for particle formation to understand how to control the physical and chemical characteristics of the spinel powder.
- Develop a thermodynamic and kinetic model to predict the behavior of the particle formation and subsequent physical and chemical characteristics of the spinel as a function of the processing variables.





Hydrothermal Synthesis

- Chemical precursors are heterogeneous slurries, gel and or homogeneous solutions, acid or base mineralizer required
- Aqueous, mixed solvent or solvothermal solution medium
- Mild to severe reaction conditions (T=25-600°C, p=1-4110 atm)
- Anhydrous oxides form in a single process step
- $P-T-H_2O$ interaction => unique phase equilibria
- Solution-mediated reaction => labile reaction kinetics relative to solid state reaction
- Controlled nucleation, growth and aging => controlled size and morphology
- Process can be inexpensive





Mechanochemical and M-H Synthesis

- Mechanical forces generate stresses that induce chemical reactions
- Inexpensive milling equipment
- Inexpensive starting materials
- Well suited for scale up (air, atomspheric pressure, room temperature)
- Solid-solid reactions
- Reactions between solid and aqueous solution (M-H synthesis)





Conventional- and Mechanochemical-Hydrothermal Methods

C-H method P = 5-20 bars

T = 50-200°C RPM = 0, 100-1400

M-H method

P = 1 atmT = 25-32°C RPM = 800-2000



Parr Autoclaves: Model 4530



Nara Micros: MIC-0

Rational Approach to Direct Crystallization of Oxides

- Compute thermodynamic equilibria as a function of the processing variables for phase of interest
- Generate equilibrium diagrams to map processing variable space for phase of interest
- Design hydrothermal experiments to test and validate computed diagrams
- Utilize processing variable space maps to explore opportunities for control of reaction and crystallization kinetics





Baseline Knowledge

- Thermodynamics
- Reaction Kinetics
- Crystallization Kinetics
- Reactor know-how





Principles Governing Crystal Growth



- Uniformity creates nucleation, growth, and ageing regimes
- Nucleation is an on/off function
- Growth is a continuous function once solubility (S>1) is exceeded in the slightest fashion
- Ageing is dominant after S approaches one (saturated solution)





Equilibria of $Ca(OH)_2$ - H_3PO_4 - NH_4OH - HNO_3 - H_2O System 1. $H_2O = H^{+1} + OH^{-1}$ 21. $CaHPO_4$ (s) = $Ca^{+2} + HPO_4^{-2}$

21. CaHPO₄ (s) = Ca⁺² + HPO₄⁻² 22. CaHPO_{4 2}•H₂O (s) = Ca⁺² + HPO₄-²+ 2H₂O 23. $Ca_3(PO_4)_2$ (s) = $3Ca^{+2} + 2PO_4^{-3}$ 24. $Ca(H_2PO_4)_2 \bullet H_2O(s) = Ca^{+2} + 2H_2PO_4^{-1} + H_2O$ 25. $Ca(H_2PO_4)_2$ (s) = $Ca^{+2} + 2H_2PO_4^{-1}$ 26. $Ca_{4}O(PO_{4})_{2}(s) + H_{2}O = 4Ca^{+2} + 2OH^{-1} + 2PO_{4}^{-3}$ 27. $Ca_{10}O(PO_4)_6$ (s) + H₂O = 10Ca⁺² + 2OH⁻¹ + 6PO₄⁻³ 28. $Ca_{4}H(PO_{4})_{3}$ (s) = $4Ca^{+2} + HPO_{4}^{-2} + 2PO_{4}^{-3}$ 29. $Ca_8H_2(PO_4)_{6.5} \bullet H_2O(s) = 8Ca^{+2} + 2HPO_4^{-2} + 4PO_4^{-3} + 5H_2O$ 30. $Ca(NO_3)_{2,3}H_2O(s) = Ca^{+2} + 2NO_3^{-1} + 3H_2O$ 31. $Ca(NO_3)_{24}H_2O(s) = Ca^{+2} + 2NO_3^{-1} + 4H_2O$ 32. $Ca(NO_3)_2(s) = Ca^{+2} + 2NO_3^{-1}$ 33. $Ca(OH)_2$ (s) = $Ca^{+2} + 2OH^{-1}$ 34. $(NH_4)_2HPO_4.2H_2O(s) = 2NH_4^{+1} + HPO_4^{-2} + 2H_2O$ 35. $(NH_{4})_{2}HPO_{4}$ (s) = $2NH_{4}^{+1} + HPO_{4}^{-2}$ 36. $(NH_4)_3PO_{43} \bullet 3H_2O(s) = 3NH_4^{+1} + PO_4^{-3} + 3H_2O$ 37. $(NH_{4})H_{2}PO_{4}$ (s) = $NH_{4}^{+1} + H_{2}PO_{4}^{-1}$ 38. $(NH_4)NO_3$ (s) = $NH_4^{+1} + NO_3^{-1}$ 39. $H_2O(v) = H_2O$ 40. $NH_3(v) = NH_{2}(aq)$ 41. HNO_3 (v) = HNO_3 (aq)

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2. $HP_2O_7^{-3} = H^{+1} + P_2O_7^{-4}$

6. $HPO_{4}^{-2} = H^{+1} + PO_{4}^{-3}$

7. $H_2PO_4^{-1} = H^{+1} + HPO_4^{-2}$

8. $2 H_2 PO_4^{-1} = (H_2 PO_4)_2^{-2}$

10. HNO₃ (aq) = $H^{+1} + NO_3^{-1}$

9. $H_3PO_4(aq) = H^{+1} + H_2PO_4^{-1}$

11. $NH_3(aq) + H_2O = NH_4^{+1} + OH^{-1}$

12. NH_4NO_3 (aq) = $NH_4^{+1} + NO_3^{-1}$

13. $CaH_2PO_4^{+1} = Ca^{+2} + H_2PO_4^{-1}$

17. $CaHPO_4$ (aq) = $Ca^{+2} + HPO_4^{-2}$

18. $Ca(OH)_{2}$ (aq) = Ca⁺² + 2OH⁻¹

19. $Ca(NO_3)_2(aq) = Ca^{+2} + 2NO_3^{-1}$

20. $Ca_5(OH)(PO_4)_3 s = 5Ca^{+2} + OH^{-1} + 3PO_4^{-3}$

14. $CaNO_3^{+1} = Ca^{+2} + NO_3^{-1}$

15. $CaOH^{+1} = Ca^{+2} + OH^{-1}$

16. $CaPO_{4}^{-1} = Ca^{+2} + PO_{4}^{-3}$

3. $H_2P_2O_7^{-2} = H^{+1} + HP_2O_7^{-3}$

4. $H_{3}P_{2}O_{7}^{-1} = H^{+1} + H_{2}P_{2}O_{7}^{-2}$

5. $H_4P_2O_7(aq) = H^{+1} + H_3P_2O_7^{-1}$



Ca(CH₃COO)₂-K₃PO₄-H₂O System







Reaction Conditions

Morphology	Ca/P	NH₃/KOH	EDTA/Ca	Time (h)	Temp (°C)	Stirring Rate (rpm)
Plates	2	$\rm NH_3$	0	8	180	300 (High Shear)
Needles	2	None	0	24	200	0
Equiaxed Spheres	2	$\rm NH_3$	0	24	200	1700
Equiaxed Hexagons	1.24	КОН	1	24	200	0
Barrels	1.24	КОН	1	1	200-230	5375 (High Shear)
Dendrites	1.24	кон	1	25	192	0
Coral	2	кон	0	24	180	0
Leaves*	1.67	NH_3	0.93	24	120	0

*solvothermal: 1,4-butandiol/H₂O=30/10 (v/v)





Gallery of Morphologies with Solution Crystallization of HA





TEM – Very Small Nanoparticles! Very Difficult to Isolate.







Method of Attack

- Daniel Kopp (1st year graduate student)
 - BS Materials Science and Engineering, Rutgers University
- Thermodynamic modeling
 - Precursor selection
 - Reaction conditions
- Down-select Synthesis Approach
 - Hydrothermal synthesis
 - Conventional Reactors
 - Microwave reactors (largest one in the USA in a university)
 - Mechanochemical synthesis
 - Unique Micros 0 system at RU (only 2 in the USA)
 - Mechanochemical-hydrothermal
- Develop in situ characterization methods
 - FT-IR spectroscopy
 - XRD
- Study reaction and crystallization kinetics to control physical and chemical characteristics.
- Develop kinetic model to semi-quantitatively predict the physical and chemical characteristics of the spinel powder.
- Supply powder to Haber group in 20-50 g quantities for processing studies





Prior work - Atakan and Riman 2007

- Recommended precursors:
 MgO and Al₂O₃
- $T = 360 \,^{\circ}C$
- P = 2828 psi
- pH = 5.8
- Advantages:
 - Does not require mineralizer to control pH
 - No by-product
 - Minimal corrosion
 - Precursors available within the desired purity level
 - Are synthesis routes with cheaper precursors such as bauxite compounds feasible?

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Prior work – Atakan and Riman 2007

- Spinel hydrothermally synthesized from steam-based reactions of MgO and Al₂O₃ at 400 °C, 2680 psi and washed in 1 M HCI
- Simple separation technology!



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Mechanochemical Synthesis

- Domanski et. al.¹ synthesized spinel from MgO and Alumina at room temperature by ball milling in 140 h.
- Contamination due to milling media equipment.
- Suchanek et. al.² synthesized hydroxyapatite at room temperature using stabilized zirconia media and liners in a mechanochemical-hydrothermal (M-H) reactor.
- Can *M-H* be used to synthesize spinel, with or without hydrothermal solutions?



Nara Machinery Co.'s MIC 0 zirconia ring/zirconia-lined wet milling machine from the Riman laboratory.

- 1. J. Am. Ceram. Soc., 87 (11): 2020-2024 2004
- 2. Biomaterials, 25(19): 4647-4657 2004.





In situ characterization

- ATR-FTIR spectroscopy reaction probe technology under development in the Riman laboratory for hydrothermal processes.
- Use of beryllium cell-based Xray diffraction at PNNL, BNL and eventually at Rutgers.







Cost Function for MgO Powder

Price function for MgO powder







Cost Function for γ -Al₂O₃ Powder



99.99% pure, surface area 70-100 m²/g





Project Benefits

- Economic
 - Capital equipment, energy and raw materials costs (< \$20/ton) are kept low because of minimal drying and elimination of milling, separation and high temperature processing steps.
 - New source of $MgAl_2O_4$ spinel powder.
 - Opportunity for commercialization
- Academic
 - First low temperature spinel synthesis method that also offers control particle size and morphology.
 - First fundamental study on solution spinel crystallization thermodynamics and kinetics.
 - Method may be applicable to other ceramic chemistries





Next Steps

- Complete literature review of spinel synthesis literature and write a review paper.
- Prepare and characterize initial spinel samples using the Atakan and Riman approach.
- Research *in situ* and *ex situ* characterization techniques.
- Run and validate thermodynamic models for different precursors.
- Get training on hydrothermal reactor and chemical and physical characterization instrumentation



