Materials 218/Chemistry 277: Take home final

Date: March 13, 2019; Due date: March 19, 2019 (slip under Ram's door by 5:00 pm)

Please properly attribute the source of any material you might make use of (URLs, references ...) Also, do please read this exam sheet very carefully!

Important References

- 1. R. Hoffman and C. Zheng, J. Phys. Chem. 89 (1985) 4175-4181. [doi]
- 2. S. Jia, S. Chi, J. W. Lynn, and R. J. Cava, Phys. Rev. B 81 (2010) 214446. [doi]
- 3. S. Jia, P. Jiramongkolchai, M. R. Suchomel, B. H. Toby, J. G. Checkelsky, N. P. Ong, and R. J. Cava, *Nature Phys.* 7 (2011) 207–210. [doi]

Questions

1. Fascinating properties have been discovered in compounds with the ThCr₂Si₂ structure type ("122" structure) with the generic formula, AM_2X_2 . Some of these compounds have $A = Ca^{2+}$, X = P, and M is a (formally divalent) 3d transition metal (*i.e.* Fe²⁺, Co²⁺, Ni²⁺). Sketch the structures of CaFe₂P₂, CaCo₂P₂, and CaNi₂P₂ in VESTA: Space group I4/mmm (#139), Ca = (0,0,0); B = (0,0.5,0.25); P = (0,0,z) using the data tabulated below. Is there a centering operation in this space group?

Compound	a (Å)	c (Å)	z (P)
CaFe ₂ P ₂	3.855	9.985	0.3643
$CaCo_2P_2$	3.858	9.593	0.3721
$CaNi_2P_2$	3.916	9.363	0.3774

A. Mewis, Z. Naturforsch. B 35 (1980) 141-145.

- 2. What are the coordination geometries of the *M* and P atoms (coordination numbers, *M*–P–*M* and P–*M*–P bond angles, bond lengths)? What is the shortest interlayer P–P distance in the three compounds?
- 3. Draw the electrostatic valence net for $CaFe_2P_2$ assuming the formal valences of Ca^{2+} , Fe^{2+} , and P^{3-} . Do the charges balance?
- 4. What are the bond valence sums for Ca and Fe in CaFe₂P₂? Can you infer anything about the nature of bonding in, and between the layers? Use *B* = 0.37, and *R*₀ = 2.27 Å for Fe–P and *R*₀ = 2.55 Å for Ca–P bonds.
- 5. Below are the calculated densities of states for CaFe₂P₂, CaCo₂P₂, and CaNi₂P₂. Label regions with major contributions from the *M*-*d* (*t*₂ and *e* levels), and P-*p* orbitals. [5]
- 6. Would you expect metallic conductivity in any of these compounds? For the metallic compounds, where do the metallic states come from (*M* or P)? Why? What does this tell you about the nature of bonding of the metal atoms in the layers? [5]

[4]

[4]

[4]



- 7. Which compound is likely to be magnetic in light of the Stoner criterion for itinerant magnetism and how can you tell?
- 8. Read the paper by Hoffman and Cheng (Ref. [1]). In their analysis of the stacking of $[Mn_2P_2]^{2-}$ layers, they observe strongly localized P p_z orbitals projecting into the interlayer space. According to the authors, why do the P–P distances contract as one moves across the periodic table (M = Fe to Co to Ni)? Explain with a sketch of the energies of the P p_z bonding/antibonding orbitals relative to the d bands and the Fermi level.
- 9. How would the substitution of Ge for P change the X-X distance (*e.g.* $SrCo_2(P_{1-x}Ge_x)_2$)? Discuss in the context of Hoffman & Cheng. Describe the conclusion drawn by S. Jia *et al.* (2011) in Ref. [3] for $SrCo_2(P_{0.5}Ge_{0.5})_2$ with a sketch of the DOS for the $X-X \sigma^*$ and Co 3*d* orbitals relative to the Fermi energy. Reminder: chemical potential $\approx E_F$. Would *you* expect this compound to be magnetic in light of the Stoner criterion?

[2]

[5]