## Computational Chemistry Spring Semester 2010 ( Due 5 / 3 / 10 )

- 1. Consult Mochida, K.; Matsuhida, N.; Sato, R.; Nakadaira, Y. *Organometallics* **2006**, *25*, 4231.
  - a. Scheme 2 (page 4232) shows the reaction for the thermolysis of 1 to benzene and singlet dimethylgermylene, Ge(Me)<sub>2</sub>, and suggests a possible transition state (3). Determine whether 3 is really a transition state or whether it is a reactive intermediate (i.e., a high-energy minimum). Symmetry might prove useful in speeding up your calculations. Use M06-2X/6-31G(d) for your final assessment, although initial calculations at a lower level of theory and/or with a smaller basis set may prove useful at your discretion. Is 3 a TS or a minimum? If 3 is a minimum, find the transition-state structure between 1 and 3, and the TS structure for dissociation. Nota bene: when imposing symmetry, one occasionally finds a TS structure associated with a molecular motion that breaks the symmetry, but that is not the reaction coordinate. Visualize your imaginary modes to decide if any TS structure is actually the one that you want.
  - b. Using whatever transition state you determine to be relevant, compare your calculated  $\Delta H^\ddagger$  and  $\Delta S^\ddagger$  to the experimental values given in the paper (assume the temperature difference between default 298 K in Gaussian and the experiment to have a negligible effect on the computed thermodynamic variables). Use your predicted  $\Delta G^\ddagger$  to calculate the unimolecular rate constant, k, at 305.3 K (indicate the equation that you used to get the rate constant). Do your errors seem reasonable given the method and basis set used? Correct your electronic energies by doing *single point* calculations at the M06-2X/6-311+G(2df,p) level. Are things improved by comparison to experiment? While it is always possible that deviations between theory and experiment represent inaccuracy in the modeling, what *physical* effects might be responsible for differences between theory and experiment?
  - c. In the paper, the progress of the reaction is described as having been followed by proton NMR, but the shifts that were observed are not described. Could the reaction have been followed instead using UV/Vis spectroscopy? Based on TD DFT computed UV/Vis data (ask for 10 states in your calculations (do not use the default value of 3) and employ the PBE0 functional, which in G09 is invoked by the keyword pbe1pbe), how would you expect the UV/Vis spectrum to evolve over the course of the reaction?
- 2. Here continues a problem that will carry over to the final exam. We add to the data at:

Your present task is to update your two structures on the potential energy surface (PES) with a calculation at the M06/6-31G(d) level (since there is no double slash, this implies re-optimizing the geometry). The most efficient way to do this will be to read in the force constants and geometry from your RHF frequency calculations (i.e., fopt=(readfc) geom=checkpoint, guess=read) as keywords. *Nota bene*: if you were using fopt=ts you will need to continue to use ts, of course. Once you have a reoptimized structure, do a new frequency calculation at the M06/6-31G(d) level and update your entry with those data.

Finally, compute the M06/6-31G(d) *aqueous* solvation free energy for your DFT structure using the SMD model in Gaussian09 (as a *single-point* calculation on the DFT optimized geometry). When the calculation completes, add the solvation free energy to the website data. Report only one position after the decimal place for the solvation free energy in units of kcal/mol (the solvation free energy is the difference between the M06 SCF energies in solution (found in the SMD output file) and in the gas phase (found in the M06 geometry optimization output file)).