References

- Damon M. R., Kameyama M. C., Knox M. R., Porter D. H., Yuen D. A., Sevre E. Interactive visualization of 3D mantle convection // Visual Geosciences. — 2008. — P. 49—57. — DOI: 10.1007/s 10069-007-0008-1.
- Greensky J., Czech W. W., Yuen D. A., Knox M., Damon M. R., Chen S. S., Kameyama M. C. Ubiquitous interactive visualization of 3D mantle convection using a web portal with java and AJAX framework / Visual Geosciences. 2008. P. 105—115. DOI: 10.1007/ s10069-008-0013-z.
- McLane J., Czech W. W., Yuen D. A., Knox M., Wang S. M.
 Ubiquitous Interactive Visualization of Large-Scale
 Simulations in Geosciences Over a Java-based
 Web-Portal. Concurrency and Computation:
 Practice and Experiences. 2009. (in press).

- Porter D. H. Volume visualization of high-resolution data using PC Clusters. — 2002. — http:// www.lcse.umn.edu/hvr/pc_vol/rend_Lpdf/.
- Porter D. H., Woodward P. R., Iyer A. Initial experiences with grid-based volume visualization of fluid low simulations on PC Clusters. 2002. http://www.lcse.umn.edu/dhp1/ articles.html.
- Woodward P. R., Porter D. H., Greensky J., Larson A. J., Knox M., Hanson J., Ravindran N., Fuchs T. Interactive volume visualization of fluid flow simulation data. Applied Parallel Computing, State of the Art in Scientific Computing, Proc. PARA 2006 Conf., LNCS 4699, Heidelberg: Springer Verlag, 2007. P. 659—664. http://www.lcse.umn.edu/para06.

Molecular Paleoclimatology: Quantum Chemistry and the History of the Earth's Atmosphere

© J. Rustad, 2010

University of California, Davis, CA USA james.rustad@gmail.com

The history of the composition of the Earth's atmosphere is one of the most important problems of our time. As the human race considers the possibility of planetary engineering to mitigate the potential effects of CO₂-induced global warming, clearer insight into the history of atmospheric PCO2 and global temperature is required. For example, if PCO₂ values were 20 times higher at times in the geologic past, it may be premature to worry about anthropogenic increases in PCO₂ a factor of two above Pleistocene-Holocene values, which are anomalously low from the point of view of geologic history. Two major techniques for determining the history of atmospheric PCO₂ are the chemically based. The first depends on the isotopic composition of carbon incorporated into soil minerals during their growth. The second depends on the isotopic composition of boron incorporated into marine minerals during precipitation. Both methods depend crucially on knowing the equilibrium constant for isotope exchange between minerals, aqueous, and gas phases. There are major uncertainties in these equilibrium constants which strongly affect geochemical estimation of the history of PCO2 in Earth's atmo-

sphere. Here, I describe how quantum chemistry techniques can be used to reduce this uncertainty and make more reliable estimates of PCO₂ values.

Ab initio molecular dynamics and quantum chemistry techniques are used to calculate the structure, vibrational frequencies, and carbon-isotope fractionation factors of the carbon dioxide component [CO₂(m)] of soil (oxy)hydroxide minerals goethite, diaspore, and gibbsite. We have identified two possible pathways of incorporation of CO₂(m) into (oxy)hydroxide crystal structures: one in which the C4+ substitutes for four H+ [CO2(m)A] and another in which C⁴⁺ substitutes for (Al³⁺,Fe³⁺)+H⁺ [CO₂(m)_B]. Calculations of isotope fractionation factors give large differences between the two structures, with the $CO_2(m)_A$ being isotopically lighter than $CO_2(m)_B$ by≈10 per mil in the case of gibbsite and nearly 20 per mil in the case of goethite. The reduced partition function ratio of CO₂(m)_B structure in goethite differs from CO₂(g) by<1 per mil. The predicted fractionation for gibbsite is>10 per mil higher, close to those measured for calcite and aragonite. The surprisingly large difference in the carbon-isotope fractionation factor between the CO₂(m)_A and CO₂(m)_B structures within a given mineral suggests that the isotopic signatures of soil (oxy) hydroxide could be heterogeneous.

Density functional and correlated molecular orbital calculations (MP2) are carried out on B(OH)₃·nH₂O clusters (n = 0, 6, 32) and B(OH)₄nH₂O (n = 0, 8, 11, 32) to estimate the equilibrium distribution of ¹⁰B and ¹¹B isotopes between boric acid and borate in aqueous solution. For the large 32-water clusters, multiple conformations are generated from ab initio molecular dynamics simulations to account for the effect of solvent fluctuations on

the isotopic fractionation. We provide an extrapolated value of the equilibrium constant \acute{a}_{34} for the isotope exchange reaction $^{10}B(OH)_3(aq) + ^{10}B(OH)_4(aq) = ^{11}B(OH)_3(aq) + ^{11}B(OH)_4$ (aq) of 1.026—1.028 near the MP2 complete basis set limit with 32 explicit waters of solvation. With some exchange-correlation functionals we find potentially important contributions from a tetrahedral neutral $B(OH)_3 \cdot H_2O$ Lewis acid—base complex. The extrapolations presented here suggest that DFT calculations give a value for $103 \ln \acute{a}_{34}$ about 15 % higher than the MP2 calculations.

Interaction of earthquakes and slow slip: Insights from fault models governed by lab-derived friction laws

© N. Lapusta, 2010

California Institute of Technology, Pasadena, CA, USA

Motion of plates in the Earth crust is accomodated through fault slip. That includes both fast events (earthquakes) and slow relative motion, as evidenced by seismic and geodetic observations. We study mechanics and physics of earthquakes using a unique simulation approach that reproduces both earthquakes and slow slip, with full inclusion of inertial effects during simulated earthquakes, in the context of a 3D fault model. The approach incorporates laboratory-derived rate and state friction laws, including the effects of

shear heating during rapid, seismic slip, involves slow, tectonic-like loading, resolves all stages of seismic and aseismic slip, and results in realistic rupture speeds, slip velocities, and stress drops. Our simulations show that a number of observed earthquake phenomena can be explained by interaction of earthquakes and slow slip, including transition to intersonic rupture speeds during earthquakes, peculiar properties of small repeating earthquakes, and complex spatiotemporal patterns of earthquake sequences.