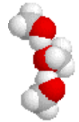
Emerging Methods in Computational Chemistry and Materials Science

Aberdeen MD, April 29-30, 1999

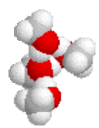
How to use standard electronic structure tools for the theory of liquids: Quasi-Chemical Theories of Associated Liquids



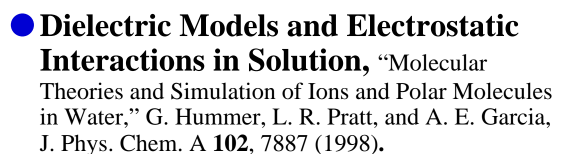
R. L. Martin, P. J. Hay, L. R. Pratt, and R. A. LaViolette



URL - http://www.t12.lanl.gov/~www/projects/ solvents.html



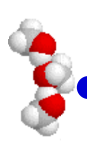
Dielectric Models -> **Molecular Theory**





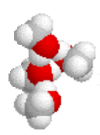
"Hydrolysis of Ferric Ion in Water and Conformational Equilibrium," R.

L. Martin, P. J. Hay, and L. R. Pratt, J. Phys. Chem. A **102**, 3565 (1998).



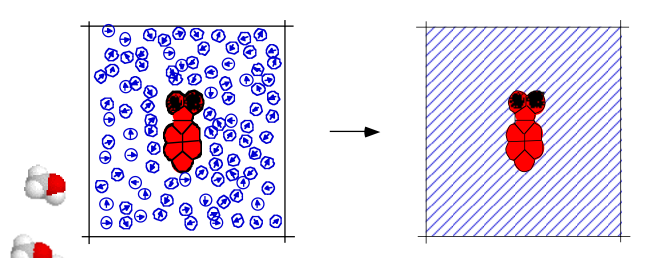
"Quasi-chemical Theories of Associated Liquids," L. R. Pratt and R. A. LaViolette, Molec. Phys. 94, 909 (1998).

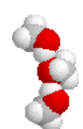
Example: liquid methanol (work in progress, with R. A. LaViolette)



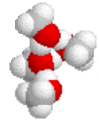
Dielectric Model

premier example of an "implicit water model"





 $\nabla \bullet \boldsymbol{\varepsilon}(\mathbf{r}) \nabla \Phi(\mathbf{r}) = -4\pi \boldsymbol{\rho}(\mathbf{r})$



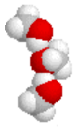
Dielectric Model



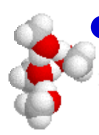
- physical can be derived from a Hamiltonian (in more than one way!)
- practical reasonably treats huge effects that can't be ignored
- chemical can simultaneously include reasonable chemistry
- empirical parameterizations (radii) are not unreasonable

con:

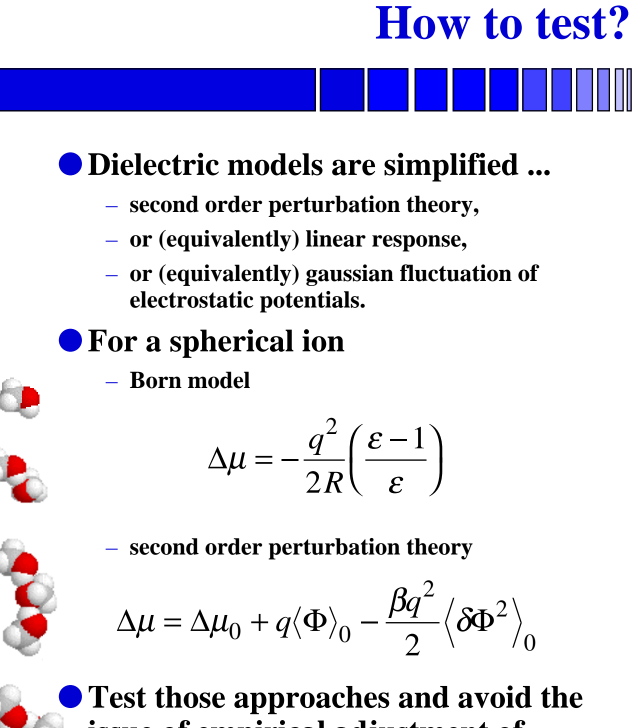
 careful, molecular scale validation of the statmech is <u>MIXED</u>!!!



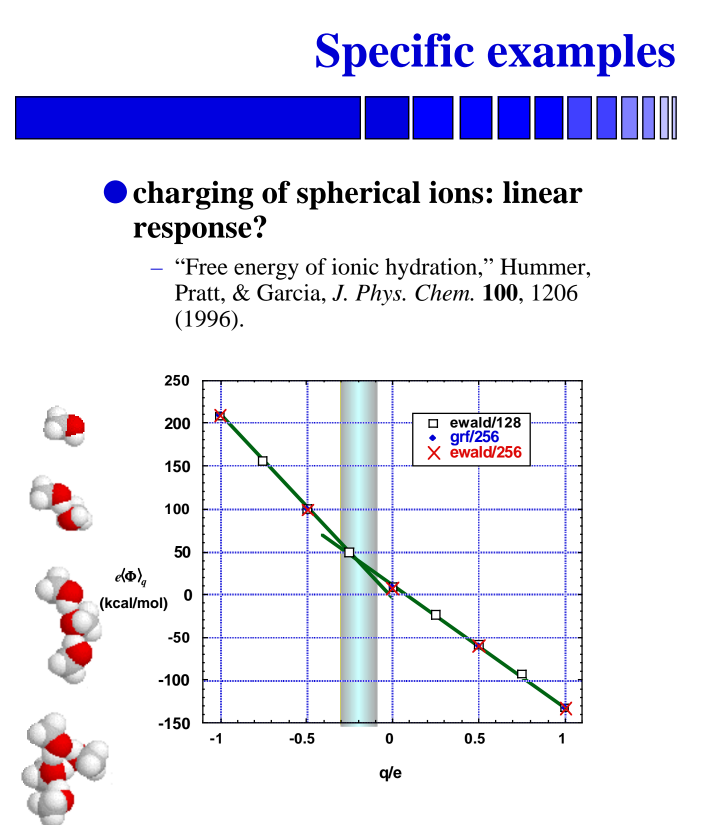
sometimes 100% empirical - parameters
 (radii) must be reconsidered for EVERY new
 result; parameters depend on (T,p,x); results
 are sensitive to parameters; any physical
 result (correct or not) can be reproduced

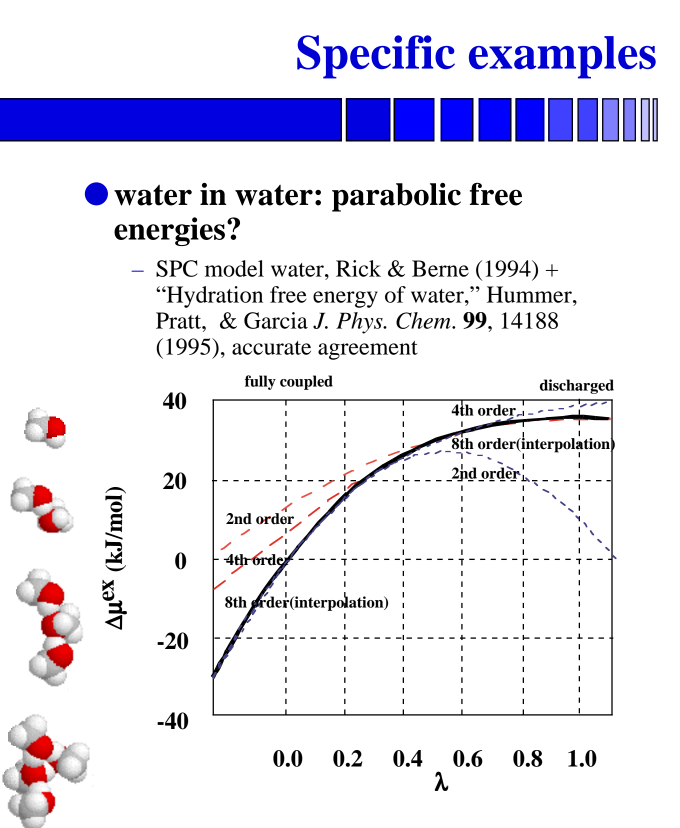


"Everything should be made as simple as possible, but no simpler." A. Einstein









Physical conclusions from simulations

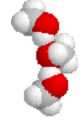
Dielectric models of hydration fail on a molecular scale when proton (H) structures near the solute lead to non-gaussian fluctuations of electrostatic potentials.

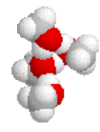
- Easiest examples to get "right" are classic cations, *e.g.* Na⁺.
- Neutral, polar, H-bonding molecules, e.g. H₂O or imidazole are more difficult cases for dielectric models when tested on a molecular scale. However, hydration free energies are smaller than for ions.

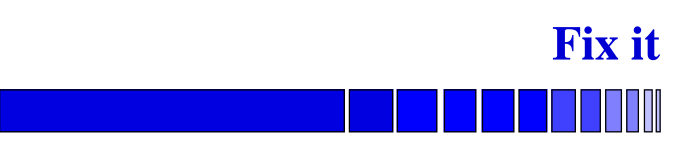
 Negative ions are a problem and the hydration free energies are again large.
Here chemistry will be more important also, *e.g.* HCO₃⁻.







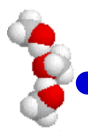




Conformational substates

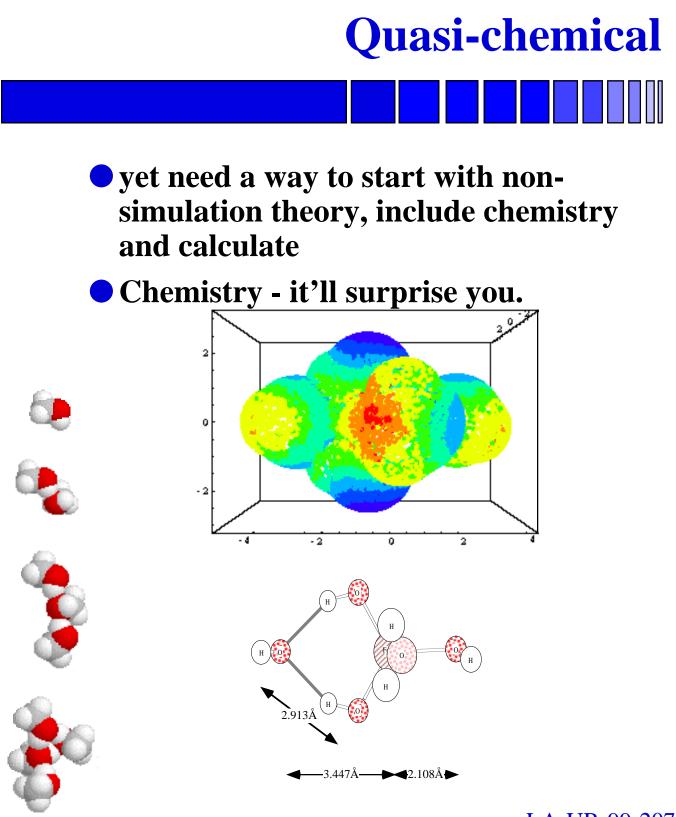
- Gaussian model for each substate. "Multistate gaussian model for electrostatic solvation free energies," Hummer, Pratt, & Garcia, J. Am. Chem. Soc. 119, 8523 (1997)
- Not "dielectric saturation and electrostriction;" incrementally higher perturbation theory not that helpful - kinks!
- Substates are categorized by numbers of close H-bonds to solute.

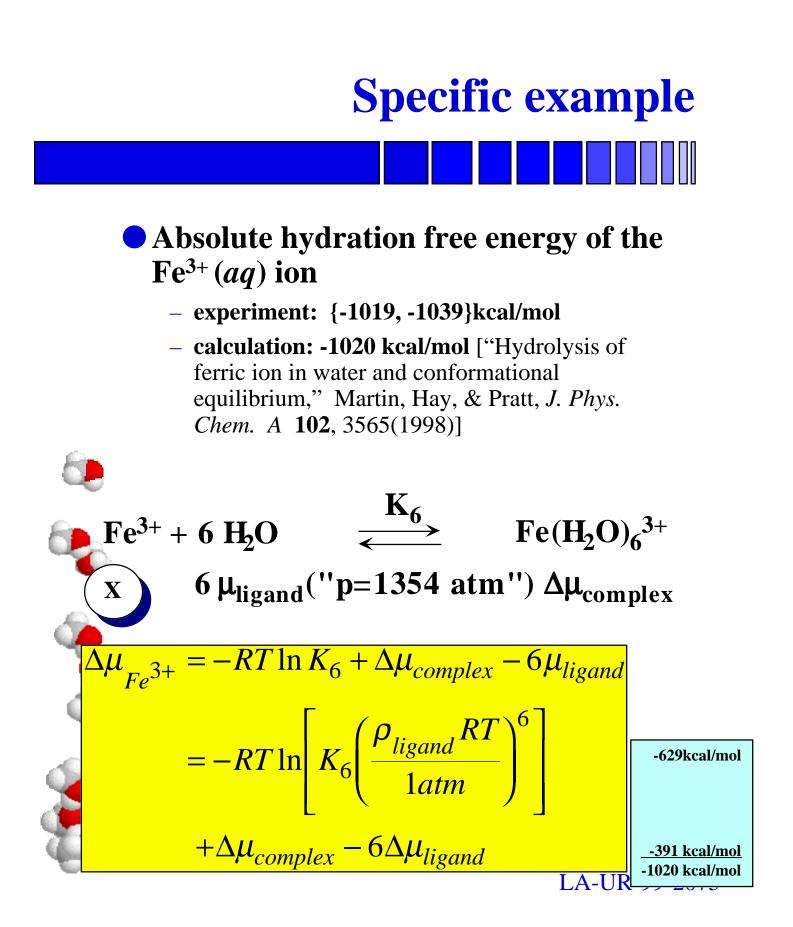
$$\beta \Delta \mu(\lambda) = -\ln \sum_{n} w_{n} e^{-\beta \lambda m_{n} + (\beta \lambda \sigma_{n})^{2}/2}$$

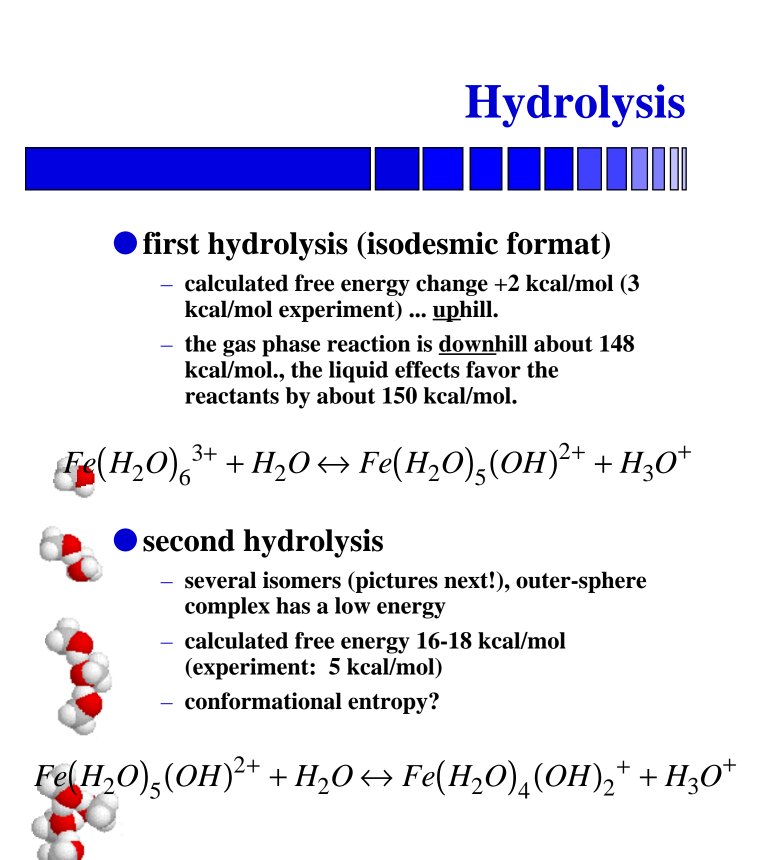


water: w_n , $1 \le n \le 6$, <n>=3.64, remaining max{error} about 5%.

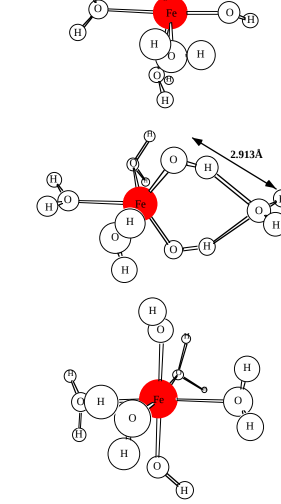
negative ions, *e.g.*, Cl⁻, still a problem.







isomeric structures for $Fe(H_2O)_4(OH)_2^+$

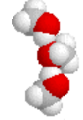


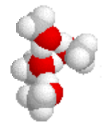
H)

Ό



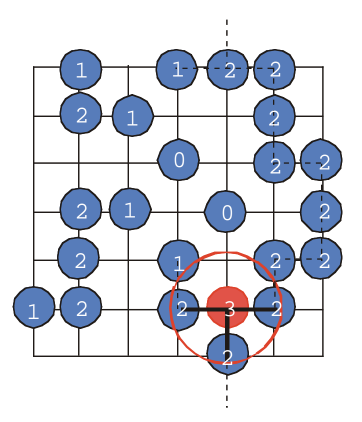






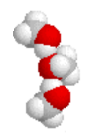
clustering...

local (not spanning) clusters







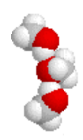


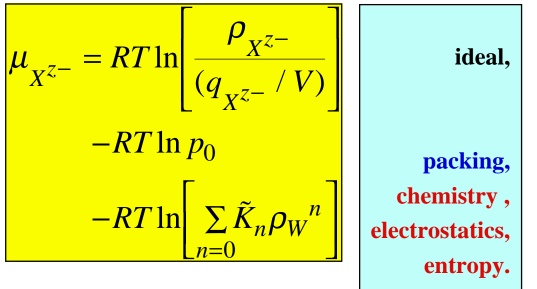
 "One of the principal objects of theoretical research in my department of knowledge is to find the point of view from which the subject appears in its greatest simplicity." J. W. Gibbs

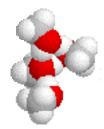
Quasi-chemical formulation

$$X^{z-} + W \underset{K_{2}/K_{1}}{\longleftrightarrow} XW^{z-}$$
$$XW^{z-} + W \underset{K_{3}/K_{2}}{\longleftrightarrow} XW_{2}^{z-}$$
$$XW_{2}^{z-} + W \underset{K_{n}/K_{n-1}}{\overset{K_{n}/K_{n-1}}{\leftrightarrow} XW_{n}^{z-}}$$
$$\vdots$$
$$XW_{n-1}^{z-} + W \underset{K_{n}/K_{n-1}}{\overset{K_{n}/K_{n-1}}{\leftrightarrow} XW_{n}^{z-}}$$



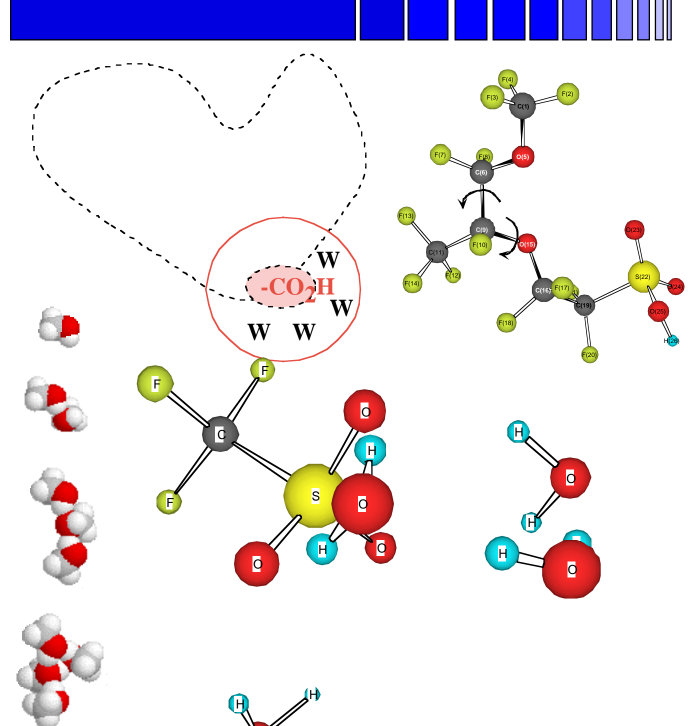




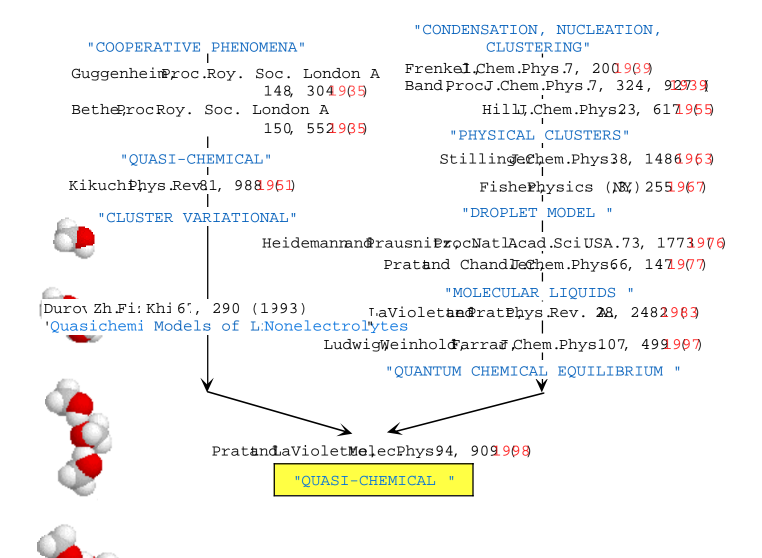


p₀=probability of an open cavity for the cluster volume, *e.g.* (1-ρv).

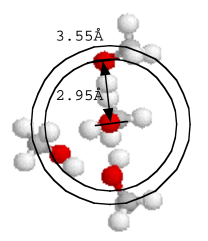
extended (macromolecules)?



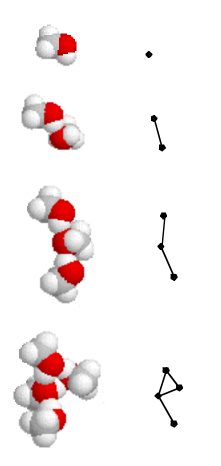


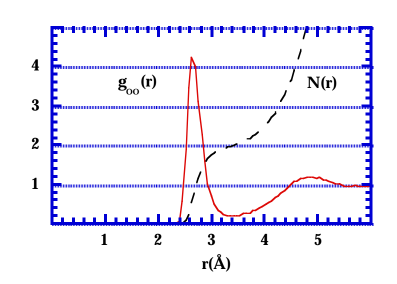


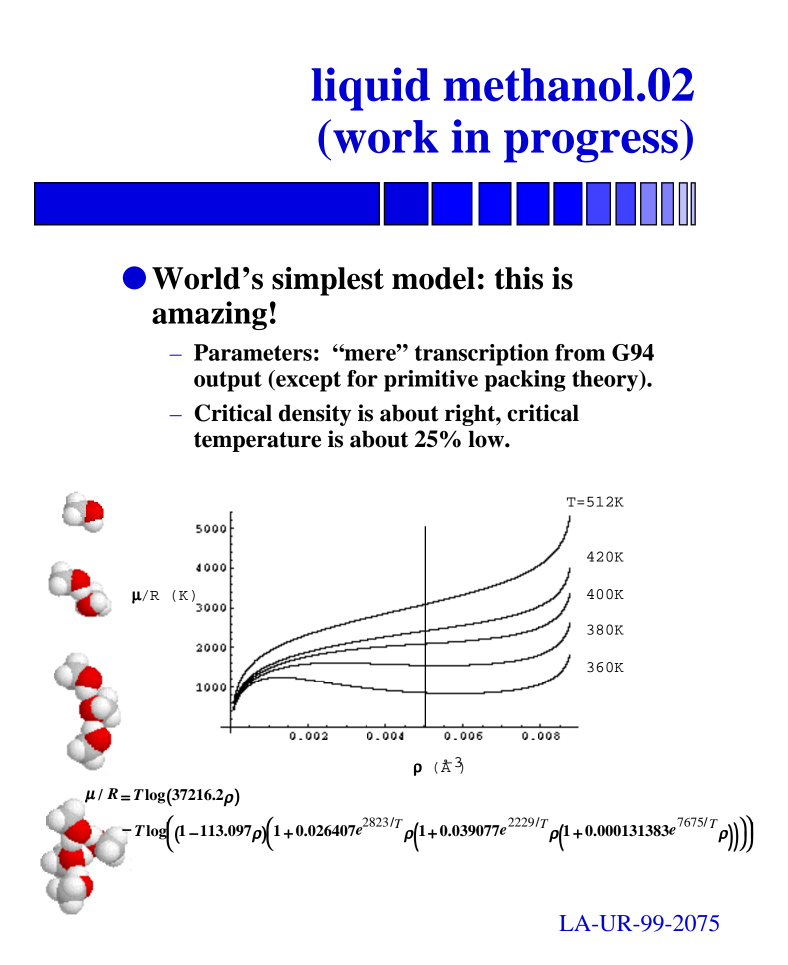
liquid methanol.01 (work in progress)

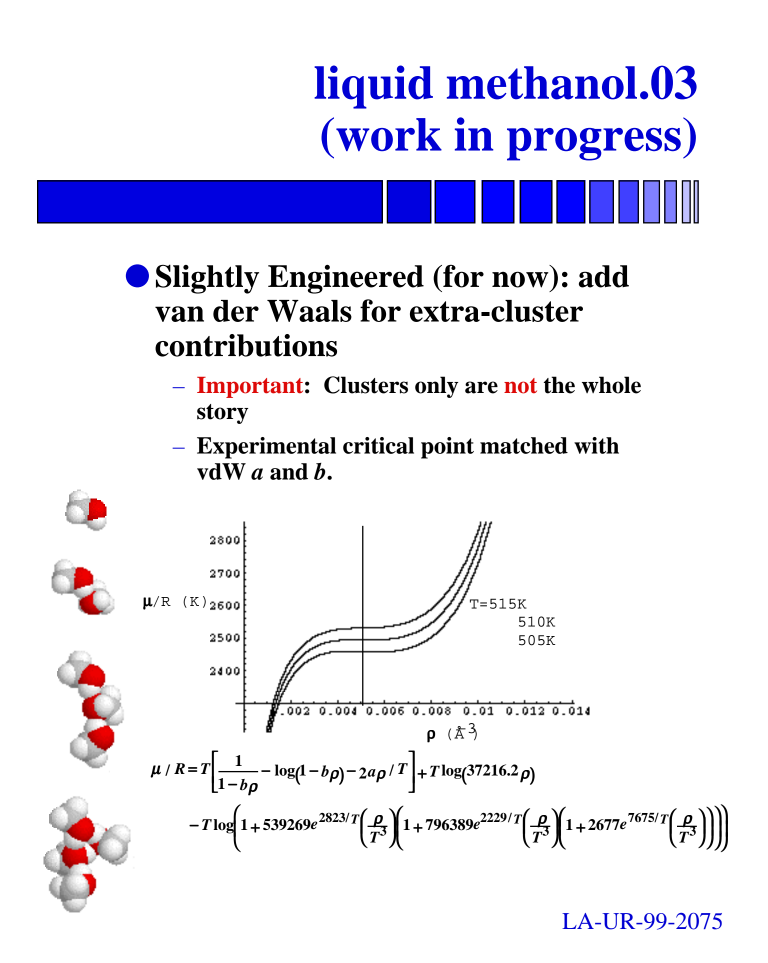


(G94, all results b3lyp/6-31g**)









liquid methanol.04 (work in progress)

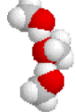
vdW parameters found are consistent with the quasi-chemical picture

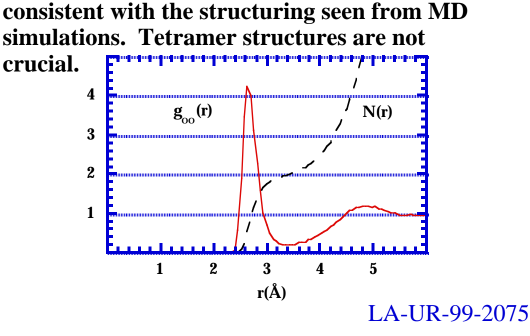
- $\rho_c a = 1.14$ kcal/mol. $2\rho a$ is the interaction potential energy of a methanol molecule with *all neighbors outside* the cluster. And this parameter value is less than typical hydrogen bonding energies. Therefore, the quasichemical theory appears to be properly and accurately isolating strong "chemical" interactions and leaving relatively weak interactions for perturbative stat-mech.

 $-\rho_c b = 0.334 \ll$ cluster radius 3.15Å. This is



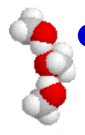






Conclusions

- Dielectric models for hydration breakdown first because of near neighbor protons that lead to nongaussian fluctuations of electrostatic potentials.
 - Quasi-chemical approaches treat inner shells specially, permit an organized application of electronic structure calculations, should also fix problems with negative ions. This is real stat-mech.



The quasi-chemical picture is clearly a "basic truth" of liquid methanol, but more work will be required to finish the stat-mech in this case.

