Spicing up continuum solvation models with SaLSA: the spherically-averaged liquid susceptibility ansatz

Ravishankar Sundararaman, Kathleen A. Schwarz, Kendra Letchworth-Weaver, and T. A. Arias
Cornell University Department of Physics, Ithaca, NY 14853, USA
(Dated: October 10, 2014)

Continuum solvation models enable electronic structure calculations of systems in liquid environments, but because of the large number of empirical parameters, they are limited to the class of systems in their fit set (typically organic molecules). Here, we derive a solvation model with no empirical parameters for the dielectric response by taking the linear response limit of a classical density functional theory for molecular liquids. This model directly incorporates the nonlocal dielectric response of the liquid using an angular momentum expansion, and with a single fit parameter for dispersion contributions it predicts solvation energies of neutral molecules with an RMS error of 1.3 kcal/mol in water and 0.8 kcal/mol in chloroform and carbon tetrachloride. We show that this model is more accurate for strongly polar and charged systems than previous solvation models because of the parameter-free electric response, and demonstrate its suitability for ab initio solvation, including self-consistent solvation in quantum Monte Carlo calculations.

Electronic density functional theory \[^{[1]}\,^{[2]}\] enables first-principles prediction of material properties at the atomic scale including structures and reaction mechanisms. Liquids play a vital role in many systems of technological and scientific interest, but the need for thermodynamic phase-space sampling complicates direct first-principles calculations. Further, absence of dispersion interactions and neglect of quantum-mechanical effects in the motion of protons limit the accuracy of ab initio molecular dynamics for solvents such as water \[^{[3]}\,^{[4]}\].

Continuum solvation models replace the effect of the solvent by the response of an empirically-determined dielectric cavity. Traditional solvation models \[^{[5]}\,^{[10]}\] employ a large number of atom-dependent parameters, are highly accurate in the class of systems to which they are fit - typically organic molecules in solution, and have been tremendously successful in the evaluation of reaction mechanisms and design of molecular catalysts. Unfortunately, the large number of parameters precludes the extrapolation of these models to systems outside their fit set, such as metallic or ionic surfaces in solution. Recent solvation models that employ an electron-density based parametrization \[^{[11]}\,^{[12]}\] require only two or three parameters and extrapolate more reliably, but still encounter difficulties for charged and highly polar systems \[^{[13]}\,^{[14]}\].

The need for empirical parameters in continuum solvation arises primarily because of the drastic simplification of the nonlocal and nonlinear response of the real liquid with that of a continuum dielectric cavity. Recently, we correlated the dielectric cavity sizes for different solvents with the extent of nonlocality of the solvent response to enable a unified electron-density parametrization for multiple solvents \[^{[15]}\], but the electron density threshold \(n_c\) that determines the cavity size still required a fit to solvation energies of organic molecules. Joint density functional theory (JDFT) \[^{[16]}\] combines a classical density functional description of the solvent with an electronic density functional description of the solute, naturally captures the nonlocal response of the fluid, and does not involve cavities that are fit to solvation energies.

Here, we derive a nonlocal continuum solvation model from the linear response limit of JDFT. Because there are no fit parameters for the electrostatic response, this theory is therefore suitable for the study of charged and strongly polar systems. This derivation begins with a simple ansatz: the distribution of solvent molecules starts out isotropic and uniform outside a region excluded by the solute; electrostatic interactions between the solute and solvent then perturb this distribution to first order. We first describe a method to estimate that initial distribution from the overlap of solute and solvent electron densities, and then show how to calculate the nonlocal linear response of the fluid using an angular momentum expansion. The addition of nonlocal cavity formation free energy and dispersion functionals derived from classical density functional theory \[^{[15]}\] results in an accurate description of solvation free energies of neutral organic molecules as well as highly polar and ionic systems. Finally, we show that the nonlocal dependence of this model on the solute electron density enables self-consistent solvation in quantum Monte Carlo calculations, which was previously impractical due to statistical noise in the local electron density \[^{[17]}\].

*Iso-density-product cavity determination* - A common ingredient in continuum solvation models is the formation of a cavity that excludes the solvent from a region of space occupied by the solute. Atom-based parametrizations typically exclude the centers of solvent molecules from a union of spheres centered on each solute atom with radius equal to the sum of the atomic and solvent van-der-Waals (vdW) radii, and then construct a dielectric cavity that is smaller by an empirical solvent-dependent radius \[^{[7]}\]. In contrast, the density-based approaches adopt a smoothly-varying dielectric constant which is a function of the local solute electron density that switches from the vacuum to bulk solvent value at a solvent-dependent critical electron density \(n_c\) \[^{[11]}\,^{[12]}\].

Here, since we explicitly account for the nonlocal response, we require only the distribution of the solvent molecule centers. Because the distance of these centers from the solute atoms corresponds to the distance of nearest approach of two non-bonded systems, vdw radii, which are defined in terms of non-bonded approach distances \[^{[19]}\], provide a reasonable guess. However, directly using the vDW radii does not account for changes in the electronic configuration between the isolated atom and molecules or solids. A description based on the electronic density would instead naturally capture this
The interaction of non-bonded systems is dominated by Pauli repulsion at short distances, which depends on the overlap of the electron densities of the two systems. Indeed, Figure 1 shows that the atom separation, $R_{12}$, at which electron density overlap $\bar{n}_c(R_{12})$ crosses a threshold value of $\bar{n}_c = 1.42 \times 10^{-3} \, \text{Å}^3$ correlates well with the sum of vdW radii $R_1 + R_2$.

Above, analogously to the Kohn-Sham approach, the liquid free energy functional employs the state of the corresponding non-interacting system specified by two sets of independent variables. The first, $p_\omega(\vec{r})$, is the orientation probability density of finding a solvent molecule centered at $\vec{r}$ with orientation $\omega \in \text{SO}(3)$. The solvent site densities $N_\alpha(\vec{r})$ are dependent variables that are calculated from $p_\omega(\vec{r})$. The second variable is the polarization density, $P_\alpha(\vec{r})$ for each solvent site.

The above analysis provides a universal threshold on the density product that can estimate the approach distance of non-bonded systems. We utilize this capability to determine the distribution of solvent molecule centers around a solute with electron density $n(\vec{r})$. Our ansatz requires a spatially-varying but isotropic initial distribution of molecules. Therefore we compute overlaps with the spherically-averaged electron density $n_{\text{eq}}^0(\vec{r}) = \int d\vec{r}_0 n_{\text{eq}}(\vec{r}_0)$, where $n_{\text{eq}}(\vec{r})$ is the electron density of a single solvent molecule and $\vec{n}$ is a unit vector. Following [14], we describe the spatial variation of the solvent distribution by the smooth ‘cavity shape’ functional

$$s(\vec{r}) = \frac{1}{2} \text{erfc} \ln \frac{n_{\text{eq}}^0(\vec{r}) * n(\vec{r})}{\bar{n}_c}$$

which smoothly transitions from vacuum ($s = 0$) to bulk fluid ($s = 1$) as the overlap of the solute and solvent densities (readily calculated as a convolution) crosses the universal overlap threshold $\bar{n}_c$.

**Nonlocal electric response** - We begin with the in-principle exact joint density-functional description [10] of the free energy of a solvated electronic system

$$A_{\text{JDFT}}[n, \{N_\alpha\}] = A_{\text{HK}}[n] + \Phi_{\text{eq}}[\{N_\alpha\}] + \Delta A[n, \{N_\alpha\}],$$

where $A_{\text{HK}}$ is the Hohenberg-Kohn functional [1] of the solute electronic density $n(\vec{r})$. $\Phi_{\text{eq}}$ is a free energy functional for the solvent in terms of nuclear site densities $\{N_\alpha(\vec{r})\}$, and $\Delta A$ captures the free energy of interaction between the solute and solvent. (See [10] for details about the theoretical framework.)

We adopt the Kohn-Sham prescription [2] with an approximate exchange-correlation functional for $A_{\text{HK}}[n]$ and the polarizable ‘scalar-EOS’ free energy functional approximation [23] for $\Phi_{\text{eq}}[\{N_\alpha\}]$. We separate $\Delta A$ in (2) as the mean-field electrostatic interaction and a remainder that is dominated by electronic repulsion and dispersion interactions. We then assume that the remainder is responsible for determining the initial isotropic distribution $N_0(\vec{r}) = N_{\text{bulk}}(\vec{r})$ of solvent molecules, where $N_{\text{bulk}}$ is the bulk number density of solvent molecules and $s(\vec{r})$ is given by (1). Substituting the free energy functional from [23], we can then write (2) as
density correlation functional for dipole rotations, and the fifth term is the potential energy for molecular polarization with site susceptibilities $\chi_\alpha$. $C_{\text{rot}}$ and $C_{\text{pol}}$ parametrize correlations in the rotations and polarization respectively, and are constrained by the bulk static and optical dielectric constants. The final term is the mean-field interaction between the solute charge density $\rho_\alpha(\vec{r})$ and the induced charge density in the liquid $\rho_{\text{lq}}(\vec{r}) - \rho_{\text{lq}}^0(\vec{r})$ (where $\rho_{\text{lq}}^0$ is the charge in the initial isotropic distribution), and the self energy of that induced charge. Here,

$$\rho_{\text{lq}}(\vec{r}) = \sum_\alpha \rho_\alpha(\vec{r}) * N_\alpha(\vec{r}) - \nabla \cdot \sum_\alpha w_\alpha(\vec{r}) * N_\alpha \tilde{P}_\alpha, \quad (4)$$

where $\rho_\alpha(\vec{r})$ and $w_\alpha(\vec{r})$ respectively specify decomposition of the solvent molecule’s charge density and nonlocal susceptibility into spherical contributions at the solvent sites. Bulk experimental properties of the liquid and ab initio calculations of a single solvent molecule constrain all involved parameters. See [23] for details; the terms above are identical, except for the inclusion of solute-solvent interactions in the final electrostatic term and for the separation of the initial isotropic contributions into $\Phi_0[N_0]$.

Next, we treat the orientation-dependent pieces perturbatively by expanding $\rho_\alpha(\vec{r}) = N_0(\vec{r})(1 + \sum_{lm} x_{lnm}^l(\vec{r}) D_l^{lnm}(\omega))$, where $D_l^{lnm}(\omega)$ are the Wigner $D$-matrices (irreducible representations of SO(3)) [24]. We then expand the free energy to quadratic order in the independent variables $x_{lnm}^l(\vec{r})$ (rotation) and $P_\alpha(\vec{r})$ (polarization), formally solve the corresponding linear Euler-Lagrange equations and substitute those solutions back into (5).

The left panels show the bound charge ($+$ red, $-$ blue) and electron density (green).

For practical calculations, we rewrite the last term of \textbf{(5)} as $\int (\phi - \phi_0) \rho_{\text{rot}}/2$, where $\phi_0 = K \rho_{\text{el}}$ is the electrostatic potential in vacuum and $\phi$ is the total (mean-field) electrostatic potential which solves the modified Poisson-like equation $(\nabla^2 + 4\pi \chi) \phi = -4\pi \rho_{\text{el}}$. The $l=1$ rotational and polarization terms in $\chi \phi$ have the structure $\nabla \cdot w * N(\vec{r}) w * \nabla \phi$ which resembles the Poisson equation for an inhomogeneous dielectric, except for the convolutions that introduce the non-locality. For neutral solvent molecules, the $l=0$ term captures the interaction of the solute with a spherical charge distribution of zero net charge, and is zero except for small contributions from non-zero but negligible overlap of the solute and solvent charges. However, note that the SaLSA response easily generalizes to mixtures, and for ionic species in the solution, the $l=0$ term converts the Poisson-like equation to a Helmholtz-like equation that naturally captures the Debye-screening effects of electrolytes as in [25].

The $l>1$ terms resemble (nonlocal versions of) higher-order differential operators and capture interactions with higher-order multipoles of the solvent molecule, which decrease in magnitude with increasing $l$. We find that including terms up to $l=3$ is sufficient to converge the solvation energies to 0.1 kcal/mol.

The nonlocality of the SaLSA response allows the fluid bound charge to appear at a distance from the edge of the cavity. For example, for a water molecule in liquid water, the SaLSA cavity transitions at about the first peak of the radial distribution function $g_{\text{OO}}(r)$ of water (Figure 2), whereas the bound charge is dominant at smaller distances. In contrast, local solvation models require a smaller unphysical cavity to produce bound charge at the appropriate distance to capture the experimental solvation energy. This key difference from the local models allows a non-empirical description of the electric response in SaLSA.

At this stage, the solvated free energy \textbf{(5)} is fully specified except for the free energy of the initial configuration $\Phi_0[N_0]$, dominated by electronic repulsion, dispersion and the free energy of forming a cavity in the liquid. We set $\Phi_0[N_0] = G_{\text{cav}}[\delta] + E_{\text{disp}}[N_0]$, where $G_{\text{cav}}$ is a non-local weighted density approximation to the cavity formation free energy and $E_{\text{disp}}$ empirically accounts for dispersion and the
remaining contributions, exactly as in [15]. (See [15] for a full specification.) Briefly, $G_{\text{cav}}$ is completely constrained by bulk properties of the solvent including density, surface tension and vapor pressure, and reproduces the classical density functional and molecular dynamics predictions for the cavity formation free energy from [22] with no fit parameters. $E_{\text{disp}}$ employs a semi-empirical pair potential dispersion correction [26] which includes a scale parameter $s_0$, which we fit to solvation energies below. Note, however, that unlike previous continuum solvation models, the dominant electrostatic response includes no parameters that are fit to solvation energies.

**Solvation energies** - We implement the SaLSA solvation model in the open source density-functional software JDFTx [27], and perform calculations using norm-conserving pseudopotentials [20] at 30 $E_h$ plane-wave cut-off and the revTPSS meta-generalized-gradient exchange-correlation functional [28]. For three solvents, water, chloroform and carbon tetrachloride, we fit the sole parameter $s_0$ to minimize the RMS error in the solvation energies of a small but representative set of neutral organic molecules with a variety of functional groups and chain lengths (same set for each solvent as in [15]). Table 1 summarizes the optimum values of $s_0$ and the corresponding RMS error in solvation energies. The RMS errors of SaLSA are only slightly higher than those of the local solvation model from [15] that includes additional fit parameters for the electric response.

![Figure 3](image-url)  
**FIG. 3.** Solvation energies in water predicted by the SaLSA nonlocal solvation model for (a) organic molecules and (b) ions compared to experiment [29–31] and the linear and nonlinear solvation models from [13].

Figure 3 compares the aqueous solvation energy predictions of SaLSA with those of the linear and nonlinear local-response models from [14]. All three models perform comparably for the neutral molecule set (Figure 3(a)) used for the parameter fit, but SaLSA is substantially more accurate for inorganic ions (Figure 3(b)). In particular, the local models severely over-predict the solvation energies of small cations, and the nonlocal SaLSA model reduces the error by a factor of three for Li+ and Na+. However, SaLSA does not correct the systematic over-solvation of cations compared to anions, a known deficiency of electron-density based solvation models [13].

Finally, we turn to solvation in diffusion quantum Monte Carlo (DMC) calculations. Conventional density-based solvation models [13][14] are sensitive to the electron density in the low density regions of space ($n(\vec{r}) \sim 10^{-4} - 10^{-3} \ a_0^{-3}$). This sensitivity imposes extremely stringent restrictions on the statistical noise in the DMC electron density, rendering self-consistent solvated DMC calculations impractical. A scheme correct to first order that combines a solvated density-functional calculation with a DMC calculation in an external potential provides reasonable accuracy without calculating DMC electron densities [17]. However, full self-consistency would be particularly important for systems where density-functional approximations fail drastically.

![Figure 4](image-url)  
**FIG. 4.** Convergence in self-consistent diffusion Monte-Carlo solvation energies for (a) carbon monoxide, (b) methanol and (c) fluoride anion in water.
proximations for anions. The solvation energy at each cycle proceeds exactly as in [17] with slight differences in the details of the DMC calculation: we use Trail-Needs pseudopotentials [33, 34] with a DFT plane-wave cutoff of 70 $E_h$, and a DMC time step of 0.004 $E_h^{-1}$. Figure 4 shows that the solvated DMC energy converges to within 0.1 kcal/mol in just three self-consistency cycles. The solvation energies of methanol and carbon monoxide are essentially unchanged from the density-functional results, which suffer from strong self-interaction errors in DFT, is corrected by 3 kcal/mol. This change, although in the right direction, is small compared to the 20 kcal/mol error in the predicted SaLSA solvation energy using DFT (Figure 3(b)). The error in the solvation of anions is therefore not predominantly caused by the inaccuracy of electronic density-functional approximations for anions.

Conclusions - The linear-response limit of joint density-functional theory, combined with an electron-density overlap based estimate of the initial solvent molecule distribution, provides a nonlocal continuum solvation model with no empirical parameters for the electric response. Consequently, this ‘SaLSA’ model extrapolates more reliably from neutral organic molecules to ions and is an excellent candidate for describing highly polar and charged surfaces in solution. Further, the nonlocality of this model enables self-consistent solvation in diffusion Monte Carlo calculations. This opens up the possibility of studying systems in solution for which standard density-functional approximations fail, such as the adsorption of carbon monoxide on transition metal surfaces. Additionally, SaLSA should facilitate the development of more accurate and perhaps more empirical models that, for example, account for the charge asymmetry in the solvation of cations and anions.

This work was supported as a part of the Energy Materials Center at Cornell (EMC2), an Energy Frontier Research Center funded by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Award Number de-sc0001086.

References

[1] P. Hohenberg and W. Kohn, Phys. Rev. 136, B864 (1964).
[2] W. Kohn and L. Sham, Phys. Rev. 140, A1133 (1965).
[3] J. C. Grossman, E. Schwegler, E. W. Draeger, F. Gygi, and G. Galli, J. Chem. Phys. 120, 300 (2004).
[4] J. A. Morrone and R. Car, Phys. Rev. Lett. 101, 017801 (2008).
[5] A. Fortunelli and J. Tomasi, Chem. Phys. Lett. 231, 34 (1994).
[6] V. Barone, M. Cossi, and J. Tomasi, J. Chem. Phys. 107, 3210 (1997).
[7] J. Tomasi, B. Mennucci, and R. Cammi, Chem. Rev. 105, 2999 (2005).
[8] C. J. Cramer and D. G. Truhlar, J. Am. Chem. Soc. 113, 8305 (1991).
[9] A. V. Marenich, R. M. Olson, C. P. Kelly, C. J. Cramer, and D. G. Truhlar, J. Chem. Theory Computation, 2011 (2007).
[10] A. V. Marenich, C. J. Cramer, and D. G. Truhlar, J. Phys. Chem. B 113, 6378 (2009).
[11] O. Andreussi, I. Dabo, and N. Marzari, J. Chem. Phys. 136, 064102 (2012).
[12] K. Letchworth-Weaver and T. A. Arias, Phys. Rev. B 86, 075140 (2012).
[13] C. Dupont, O. Andreussi, and N. Marzari, J. Chem. Phys. 139, 214110 (2013).
[14] D. Gunceler, K. Letchworth-Weaver, R. Sundararaman, K. Schwarz, and T. Arias, Modelling Simul. Mater. Sci. Eng. 21, 074005 (2013).
[15] R. Sundararaman, D. Gunceler, and T. A. Arias, “Weighted-density functionals for cavity formation and dispersion energies in continuum solvation models,” Preprint arXiv:1407.4011.
[16] S. A. Petrosyian, J.-F. Briere, D. Roundy, and T. A. Arias, Phys. Rev. B 75, 205105 (2007).
[17] K. A. Schwarz, R. Sundararaman, K. Letchworth-Weaver, T. A. Arias, and R. G. Hennig, Phys. Rev. B 85, 201102(R) (2012).
[18] W. M. Haynes, ed., CRC Handbook of Physics and Chemistry 9th ed (Taylor and Francis, 2012).
[19] M. Mantina, A. C. Chamberlin, R. Valero, C. J. Cramer, and D. G. Truhlar, J. Phys. Chem. A 113, 5806 (2009).
[20] OPIUM, Pseudopotential generation project, http://opium.sf.net.
[21] Perdew and Zunger, Phys. Rev. B 23, 5048 (1981).
[22] J. P. Perdew, K. Burke, and M. Ernzerhof, Phys. Rev. Lett. 77, 3865 (1996).
[23] R. Sundararaman, K. Letchworth-Weaver, T. A. Arias, and T. A. Arias, J. Chem. Phys. 140, 144504 (2014).
[24] E. P. Wigner, Group theory and its application to the quantum mechanics of atomic spectra (Academic Press, New York, 1959).
[25] A. K. Soper, Chem. Phys. 258, 121 (2000).
[26] S. Grimme, J. Comput. Chem 27, 1787 (2006).
[27] R. Sundararaman, K. L.-W. D. Gunceler, and T. A. Arias, “JDFTx,” http://jdftx.sourceforge.net (2012).
[28] J. P. Perdew, A. Ruzsinszky, G. J. Csonka, L. A. Constantin, and J. Sun, Phys. Rev. Lett. 103, 026403 (2009).
[29] D. J. Tannor, B. Marten, R. Murphy, R. A. Friesner, D. Sitkoff, A. Nicholls, M. Ringnalda, W. A. Goddard, and B. Honig, J. Am. Chem. Soc. 116, 11875 (1994).
[30] M. Marten, K. Kim, C. Cortis, R. A. Friesner, R. B. Murphy, M. N. Ringnalda, D. Sitkoff, and B. Honig, J. Phys. Chem. B 100, 11775 (1996).
[31] C. F. Kelly, C. J. Cramer, and D. G. Truhlar, J. Phys. Chem. B 110, 16066 (2006).
[32] R. J. Needs, M. D. Towler, N. D. Drummond, and P. L. Rios, J. Phys.: Condens. Matter 22, 023201 (2010).
[33] J. Trail and R. Needs, J. Chem. Phys. 122, 174109 (2005).
[34] J. Trail and R. Needs, J. Chem. Phys. 122, 014112 (2005).