Asymmetric primitive-model electrolytes: Debye-Hückel theory, criticality, and energy bounds

Daniel M. Zuckerman,* Michael E. Fisher, and Stefan Bekiranov†
Institute for Physical Science and Technology, University of Maryland, College Park, Maryland 20742

(Received 13 November 2000; published 21 June 2001)

Debye-Hückel (DH) theory is extended to treat two-component size- and charge-asymmetric primitive models, focusing primarily on the 1:1 additive hard-sphere electrolyte with, say, negative ion diameters $a_{---}$ larger than the positive ion diameters $a_{++}$.

The treatment highlights the crucial importance of the charge-unbalanced “border zones” around each ion into which other ions of only one species may penetrate. Extensions of the DH approach that describe the border zones in a physically reasonable way are exact at high $T$ and low density $\rho$, and, furthermore, are also in substantial agreement with recent simulation predictions for trends in the critical parameters, $T_c$ and $\rho_c$, with increasing size asymmetry. Conversely, the simplest linear asymmetric DH description, which fails to account for physically expected behavior in the border zones at low $T$, can violate a new lower bound on the energy (which applies generally to models asymmetric in both charge and size). Other theories, including those based on the mean spherical approximation, predict trends in the critical parameters quite opposite to those established by the simulations.

DOI: 10.1103/PhysRevE.64.011206

PACS number(s): 02.70.Rr, 05.70.Jk, 64.70.Fx

I. INTRODUCTION

While the values of the critical parameters for the fully symmetric hard-sphere model of an electrolyte (the “restricted primitive model” or RPM [1–3]) have received extensive theoretical attention for a number of years (see, e.g., [3–10]), only recently has interest focused on the effects of asymmetry on phase coexistence and criticality [11–17]. Sabir, Bhuiyan, and Outhwaite [11] used the mean spherical approximation (MSA) and two different Poisson-Boltzmann approaches to compute critical parameters resulting from both size and charge asymmetry; the reduced critical density $\rho_c^*$ was always reported to increase with greater size asymmetry, but the trends for the reduced critical temperature $T_c^*$ were inconsistent. (For appropriate definitions of reduced temperature and density, see [14] and below.) Gonzalez-Tovar used the MSA and found, via the energy route, that both $T_c^*$ and $\rho_c^*$ increased with greater size asymmetry [12]. Most recently, Raineri, Routh, and Stell also employed the MSA, but augmented the analysis by incorporating Bjerrum-Ebeling-Grigo pairing; they likewise predicted that both $T_c^*$ and $\rho_c^*$ increase monotonically with size asymmetry [13]. Recent simulations by Romero-Enrique et al. [14] (see also [15]), however, reveal the opposite trends, namely, that critical temperature and density decrease strongly with increasing size asymmetry. Hence, as remarked in [14], our current state of even qualitative theoretical understanding appears less than adequate.

The present report therefore formulates and analyzes extensions of the Debye-Hückel (DH) approach [1] to asymmetric primitive models [17]. We may recall that the original DH theory, when supplemented by Bjerrum ion pairing [2] and dipolar-pair solvation in the ionic fluid [9], remains the most quantitatively successful theory of coexistence and criticality in the size-symmetric RPM [see, e.g., [7(b),10]]. One might, thus, reasonably hope that a suitable extension of the DH and (DHb1D) [9] theories to unequal ion sizes will, at least, yield correct trends in the dependence of $T_c^*$ and $\rho_c^*$ on asymmetry.

In fact, in their original 1923 paper, Debye and Hückel [1] already claimed to have explicit results for asymmetric primitive models (see [17]). However, serious flaws in this historic formulation suggest other, more systematic extensions of symmetric DH theory appropriate for the asymmetric case [17]. We analyze these improved “asymmetric Debye-Hückel” (ADH) theories by comparing with exact series expansions and newly developed bounds, and also assess their general physical character. We find, in fact, that, contrary to Refs. [11–13], the critical parameters predicted by our modified ADH theories [17] exhibit trends in agreement with those obtained in the recent simulations [14,15].

In order to focus on the effects of size asymmetry, we confine the present discussion primarily to the two-species size-asymmetric primitive model (or SAPM), which consists of equal numbers, $N_+ = N_-$, of positive and negative ions with hard-core diameters $a_{---} > a_{++}$, and charges of equal magnitudes $q_+ = -q_- = \pm z q_0$. (Of course, the complementary case $a_{++} > a_{---}$ follows trivially by symmetry.) All material and space is assumed to possess a uniform dielectric constant $D$. For the most part, we also assume additivity of diameters,

$$a_{+-} = a = \frac{1}{2}(a_{++} + a_{--}). \quad (1.1)$$

The degree of asymmetry will be described either by the fractional deviations from $a$, namely,

$$\delta_a = (a_{\sigma\sigma} - a)/a \quad (\sigma = +, -), \quad (1.2)$$

or by the diameter ratio [14]...
\[ \lambda = a_{--} / a_{++} = (1 + \delta_+)//(1 - \delta_-), \]  

where we have used the fact that \( \delta_+ = -\delta_- \) for the additive case. Note that the general asymmetric primitive model electrolyte possesses a large—in fact, three-dimensional—parameter space: two degrees of freedom for size asymmetry (since additivity need not be assumed) and one for charge asymmetry (noting that the overall magnitude is irrelevant).

We take the total number density to be \( \rho = N/V \) with \( N = N_+ + N_- \) and \( V \) the volume. The reduced density is then appropriately defined by [14]

\[ \rho^* = (\rho_+ + \rho_-) a^3, \]  

which seems the most relevant parameter since the critical densities are typically low, \( \rho^*_c \leq 0.1 \). But we note that other workers have used alternative definitions [11,13]. The natural temperature scale for studying critically (i.e., low temperatures) in the SAPM is set by the energy of an attractive pair of ions at contact, so we take

\[ T^* = k_B T / (|q_+ q_-|/D a_{++}), \]  

It is also convenient to define the reduced electrostatic energy, also normalized by the energy of closest approach of a \((+, -)\) pair, namely,

\[ u(\rho, T) = U_N^a(\rho, T) / N (|q_+ q_-|/D a), \]  

where \( U_N^a (= U_N^{\text{ex}} - \frac{1}{2} N k_B T) \) is the overall excess energy.

The theoretical description of size-asymmetric models is, naturally, more complicated than in the symmetric case. Specifically, in considering the correlations and fluctuations in the neighborhood of a particular ion (which, following DH, we may suppose is fixed at the origin), three distinct surrounding spherical shells or zones must be accounted for. To see this, suppose first that the selected, fixed ion is positive and that the diameters are ordered in size according to \( a_{++} < a_{+<} < a_{--} \): we term this the “inner” case since the like-like diameter of the central charge is smaller than \( a_{--} \). Trivially, no ion center can enter the “interior” zone \( 0 < r < a_{++} \), where \( r \) is the radial distance from the origin; of course, this applies to the symmetric RPM as well. New to the SAPM, however, is the “border” zone (or, more precisely, the “sub-border” zone) \( a_{++} < r < a_{++} \) for this “inner” scenario, that is shown shaded in Fig. 1: this can be populated only by the centers of the smaller, i.e., positive, ions; the larger negative ions are excluded. Finally, as in the symmetric model, positive and negative ions may be present in the exterior zone \( r > a_{++} \). When a larger negative ion is chosen to be at the origin, there is a complementary “super-border” zone, \( a_{++} < r < a_{--} \), into which only positive charges can penetrate.

The presence of these “charge-unbalanced” zones in the asymmetric case turns out to play a crucial role which, in particular, means that the simple extension of the DH treatment (ADH theory), although in agreement in leading orders at high \( T \) with the exact series expansions [3,16,17], fails badly at lower temperatures. This is shown below in Sec. II after we develop the ADH formulation, which allows for the border zones. While the failure is not entirely surprising, since the general DH formulation relies on a high-temperature expansion, it does present a sharp contrast to the symmetric case. One striking consequence of the difficulties is the theory’s violation, at low temperatures, of an extension of Onsager’s lower bound for the (internal) energy of the RPM [18] that we establish in the Appendix for general primitive models that are asymmetric in both charge and size.

It is an interesting fact that analysis of the general primitive model with nonadditive diameters presents few additional problems in most cases. (See, e.g., [16] and references therein.) On the other hand, the nonadditive models are potentially useful in applications because they can represent short-range interactions beyond the pure Coulombic couplings. Thus if, for example, \( a_{++} = a_{--} = a_0 \), so that one has a size-symmetric model, but \( a_{++} > a_0 \), a moment’s consideration shows that unlike ions experience a strong short-range repulsion relative to the geometric ion size \( a_0 \). This competes with the ionic attraction and, thus, for example, increases the relative size of a Bjerrum pair and decreases its stability. Conversely, the case \( a_{++} < a_0 \) can be viewed as representing an enhanced short-range repulsion between like ions of a “true” diameter less than \( a_0 \). In reality, of course, the interactions in ionic systems deviate from pure electrostatics at short distances. Accordingly, although we mainly focus below on the additive case, we briefly indicate where necessary how to construct DH theories for the general, nonadditive model [17]. We believe this theory is of interest in its own right as well as providing a basis for more elaborate treatments. Further details of the linear asymmetric DH theory are also given in [17].

Because of the failures of the linear asymmetric DH theory, we have explored some modifications that, for modest degrees of asymmetry, ensure satisfaction of the energy bound of the Appendix and of thermal convexity requirements [10]; these criteria are also satisfied for large asymmetry in the critical and coexistence regions. The modified ADH theories accomplish this by specifically allowing for,
and compensating in a natural way, the charge imbalances induced by the existence of the border zones [17]. These theories are expounded in Sec. III and their predictions for criticality are examined in Sec. IV. It transpires, as mentioned above, that the simplest modified theories appear to be the first theoretical treatments to predict correctly (as judged by the simulations [14,15]) that both \( T^c \) and \( \rho^c \) fall when size asymmetry is introduced; however, for \( \lambda = a_{--}/a_{++} \geq 4 \), the critical density displays incorrect, nonmonotonic behavior. Finally, Sec. V presents a brief overview and some conclusions.

II. ASYMMETRIC DEBYE-HÜCKEL THEORY

A. Formulation of the theory

In this subsection we present the ADH theory. Since the cost in additional complexity is slight, we consider here (and in the Appendix) a general two-species model with point charges \( q_{\sigma} = z_{\sigma} q_0 \) (\( \sigma = +, - \)) centered in hard spheres of diameters \( a_{\sigma} \), with the collision diameters \( a_{\sigma} \), restricted only by \( a_{++} \leq a = a_{+-} = a_{-+} \). The Debye-Hückel procedure [1], interpreted generally, directs one to calculate the excess thermodynamic functions, based on approximations for the correlation functions that are then integrated via the “energy route” [19]. Specifically, defining \( \psi_\sigma \) to be the average electrostatic potential at a (fixed) ion of species \( \sigma \) due to all other ions, the reduced electrostatic energy, defined in Eq. (1.6), is given by

\[
\psi_\sigma = \frac{Da}{2(z_+-z_-)q_0} (\psi^\sigma_- - \psi^\sigma_+),
\]

where the superscripts \(<\) and \(>\) merely serve as reminders of the relative ion sizes, and also facilitate formulation of the nonadditive case. Other thermodynamic quantities, such as the free energy and pressure, follow in principle after suitable integrations and differentiations. We derive an explicit closed-form expression for the internal energy as a function of \( T \) and \( \rho \), and the second virial term for the pressure is also discussed.

We begin the calculation of the ion potentials \( \psi_\sigma(T,\rho) \) by fixing an ion of species \( \sigma \) at the origin, as in Fig. 1. The induced electrostatic potential, \( \phi(r) \), and corresponding charge density, \( \rho_q(r) \), are then related by the (exact) averaged Poisson equation, namely,

\[
\nabla^2 \langle \phi(r) \rangle_\sigma = -\frac{4\pi}{D} \langle \rho_q(r) \rangle_\sigma,
\]

where the subscript \( \sigma \) indicates that the averages are performed with a charge of species \( \sigma \) at the origin [19]. The ion potentials follow from the limit

\[
\psi_\sigma(T,\rho) = \lim_{r \to 0} \left( \phi(r) \right)_\sigma - \frac{q_\sigma}{Dr},
\]

which eliminates the self-interaction of the fixed charge at the origin.

The DH procedure then introduces two approximations for the average charge density, \( \langle \rho_q(r) \rangle_\sigma \). The well-known Poisson-Boltzmann approximation is followed by a linearization of the exponential in the Boltzmann factors, yielding

\[
\langle \rho_q(r) \rangle_\sigma \approx \sum_{\tau=-,+} q_{\tau} \rho_{\tau} \exp[-\beta q_{\tau} \langle \phi(r) \rangle_\tau],
\]

\[
\approx \sum_{\tau=-,+} q_{\tau} \rho_{\tau} [-1 - \beta q_{\tau} \langle \phi(r) \rangle_\tau],
\]

where \( \beta = 1/k_B T \). In the size-asymmetric models, the approximate charge density must be allowed to take a different form in each of the three distinct zones around the central charge. Thus, when the fixed charge is positive (\( \sigma = + \)) one has

\[
\langle \rho_q(r) \rangle_+ = q_+ \delta(r) \quad \text{for} \quad r < a_{++}, \quad (2.6)
\]

\[
\approx q_+ \rho_+ [1 - \beta q_+ \langle \phi(r) \rangle_+] \quad \text{for} \quad a_{++} < r < a, \quad (2.7)
\]

\[
= -(\kappa_D^2/4\pi) \langle \phi(r) \rangle_+ \quad \text{for} \quad r > a, \quad (2.8)
\]

where, as usual [19], overall electroneutrality has been imposed. Here, the inverse Debye length is defined in the standard way via

\[
(\kappa_D a)^2 = \frac{s_D^2}{4\pi a^2} (q_0^2 / 2 k_B T) \sum_{\sigma} \rho_{\sigma} a_{\sigma}^2.
\]

Note: (i) there is no approximation in the innermost exclusion zone \( (r < a_{++}) \); (ii) the subborder zone \( (a_{++} < r < a = a_{+-}) \) can contain none of the larger negative ions so that in Eq. (2.7), which represents the fundamental extension of the original DH theory, only \( q_+ \) appears on the right-hand side; and (iii) the exterior zone \( (r > a) \) follows the standard Debye-Hückel form as in the symmetric RPM [1,19].

The approximations (2.6)–(2.8) complete the reduction of the averaged Poisson equation (2.2) to the basic ADH equation for \( a = 1 \) ion. This may be solved for four unknown coefficients using standard electrostatic boundary conditions (continuity of \( \phi \) and \( \phi/\partial r \) at \( r = a_{++} \) and \( r = a_{+-} = a \) with \( \phi(r) \to 0 \) as \( r \to \infty \) [19], from which \( \psi_+ \) follows via Eq. (2.3). The closed forms for the electrostatic energy in the ADH approximation then result from combining the general expression for the energy (2.1) with the appropriate expressions for the \( \psi_\sigma \) potentials, which are presented below. If we introduce

\[
\theta_\sigma = \left[ |z_\sigma|/(z_+ - z_-) \right]^{1/2},
\]

one finds that the potential in the “inner” scenario, with a smaller central positive ion (indicated by the superscript \(<\)), may be written as


\[ \psi_{\sigma}(q_0; \rho; \{ z_{\sigma} \}, \{ \delta_{\sigma} \}) = \left( \frac{z_{\sigma} q_0}{D \alpha} \right) \left[ T^\omega (A_1^{<} + A_2^{<} + A_3^{<}) - B^{<} \right]^{C^{<}}, \]  

where the various contributions are

\[ A_1^{<} = (1 + x_D + x_D \delta_+) \cosh(\theta_+ x_D \delta_+) - 1, \]  

\[ A_2^{<} = - (\theta_+^2 + \theta_+ x_D + \theta_+ x_D \delta_+) \sinh(\theta_+ x_D \delta_+), \]  

\[ A_3^{<} = x_D \delta_+, \]  

\[ B^{<} = x_D \cosh(\theta_+ x_D \delta_+) - \theta_+ x_D \sinh(\theta_+ x_D \delta_+), \]  

\[ C^{<} = (1 + x_D + x_D \delta_+) \cosh(\theta_+ x_D \delta_+) \]  

\[ - \theta_+ (\theta_+^2 + x_D + x_D \delta_+) \sinh(\theta_+ x_D \delta_+). \]  

Note that \( \delta_+ \leq 0 \) for this case. The \( A_\sigma^{<} \) terms of \( \psi_{\sigma} \) have been grouped so that each vanishes individually when \( \delta_+ \rightarrow 0 \); in this limit of size symmetry, furthermore, the potential reduces to the standard DH result [1,3,19] for a size-symmetric primitive model, namely,

\[ \psi_{\sigma} = \psi_+ = -(z_{\sigma} q_0 / D \alpha) x_D / (1 + x_D). \]

When the diameter of the negative ion is less than \( \alpha \), one should simply switch the species subscripts in Eq. (2.11).

For the “outer” situation, with a larger central ion (indicated by superscript \( > \)), the corresponding result is

\[ \psi_{\sigma}(q_0; T, \rho; \{ z_{\sigma} \}, \{ \delta_{\sigma} \}) = \left( \frac{z_{\sigma} q_0}{D \alpha} \right) \left[ T^\omega (A_1^{>} + A_2^{>} + A_3^{<}) + B^{>} \right]^{C^{>}}, \]  

where the contributions are now

\[ A_1^{>} = (1 + x_D) \cosh(\theta_+ x_D \delta_-) - 1, \]  

\[ A_2^{>} = (\theta_+^2 + \theta_+ x_D) \sinh(\theta_+ x_D \delta_-), \]  

\[ A_3^{>} = - x_D \delta_-, \]  

\[ B^{>} = x_D \cosh(\theta_+ x_D \delta_-) + \theta_+ x_D \sinh(\theta_+ x_D \delta_-), \]  

\[ C^{>} = (1 + x_D) \cosh(\theta_+ x_D \delta_-) \]  

\[ + (\theta_+^2 + \theta_+ x_D) \sinh(\theta_+ x_D \delta_-). \]  

Naturally, \( \psi_{\sigma} \) also reduces to the symmetric DH result when \( \delta_- \rightarrow 0 \). (If \( a_+ > a_- \), one should simply switch the + and − subscripts.)

For future reference, notice that both \( \psi_{\sigma} \) and \( \psi_{\sigma} \) have readily calculated zero-temperature limits, namely,

\[ \psi_{\sigma}(T=0) = - \left( \frac{z_{\sigma} q_0}{D \alpha} \right) \frac{1}{1 + \delta_+} \]  

and

\[ \psi_{\sigma}(T=0) = - \frac{z_{\sigma} q_0}{D \alpha}, \]  

which are independent of density. Furthermore, small diameters \( \alpha \) are of interest; but \( \psi_{\sigma} \) diverges for “point” ions, i.e., in the limit \( a_{\sigma} \rightarrow 0 \) (or \( \delta_+ \rightleftharpoons 1 \)). This, in fact, serves as a warning sign, as will be seen below.

### B. Assessment of ADH theory

We now examine the predictions of the linear ADH theory. Following Debye and Hückel [1,19], the free energy is to be obtained from the internal energy in Eq. (2.1) by employing the Debye charging process (which is equivalent to using the standard thermodynamic relation for the free energy in terms of the energy). This yields the reduced excess free energy density as

\[ f^{ex}(T, \rho; \{ z_{\sigma} \}, \{ \delta_{\sigma} \}) = - A_N^{ex}(V, T)/V k_B T, \]

where \( A_N^{ex} \) is the total excess Helmholtz free energy. It must be recalled that in addition to the explicit dependence on \( q_0 \) (and, hence, on \( z \)) entering Eq. (2.1) via Eqs. (2.11) and (2.17), an implicit dependence occurs via \( \kappa_D^2 q_0 \) that enters Eqs. (2.12)–(2.16) and (2.18)–(2.22).

Except in the standard symmetric case (\( \lambda = 0 \), \( \delta_+ = 0 \) most of the integrals involved in Eq. (2.24) seem intractable. Nevertheless, one may derive an expansion for the free energy, and thence for all other thermodynamic quantities, such as the pressure, by expanding the expressions (2.11)–(2.22) and integrating term by term. By this route, we can check the theory against exactly known expansions (see e.g., [3,16,17]).

Thus, consider the low-density/high-temperature expansion for the general primitive model (with arbitrary charges \( q_{\sigma} = z_{\sigma} q_0 \) and diameters \( a_{\sigma} \) which is known to overall order \( \rho^{3/2} \) in the density. On specializing to the two-component case, this may be written as

\[ f(T, \rho) = \kappa_D^3 / 12 \pi + B_{2,n} \rho^2 \ln(\kappa_D a)/(T \beta)^{3/2} + \rho^2 \sum_{n=0}^{\infty} B_{2,n} / T^{3/2} \]

\[ + O(\rho^{5/2} \ln(\kappa_D a)), \]

where we again recall \( \kappa_D a = x_D \sqrt{\rho} \). The leading term here varies as \( \rho^{3/2} \) and represents the DH limiting law reproduced by all sensible approximations. The coefficient \( B_{2,0} \) derives from the second virial coefficient for a pure hard-sphere gas. At the present level of approximation, this may be accounted for precisely by including the hard-core (HC) free energy density, \( f^{HC} \), in the basic approximation [9]. The remaining coefficients \( B_{2,1}, B_{2,2}, B_{2,3}, B_{2,4}, \ldots \) arise from the electrostatic interactions.

Now, DH-based approximations (in common with the MSA, etc.) cannot generate the \( \rho^2 (\ln \rho)/T^3 \) term with coef-
It is, indeed, striking that the exact leading-order dependences in powers of $1/T$ are reproduced for arbitrary diameters, $a_{++}$, $a_{+-}$, and $a_{--}$ (including nonadditivity and point charges). However, one probably should not be so surprised since the arguments leading to the basic ADH equations, Eq. (2.2) with Eqs. (2.4) and (2.5) are, on reflection, clearly valid to leading orders in $\rho$ and $1/T$ even if it is not obvious how far the validity will go.

To study the ADH theory more quantitatively, we focus on the energy for the additive 1:1 case ($z_+ = -z_- = 1$). No integrations are then required. Figure 2 displays energy isochores as functions of $T^*$ at high and low temperatures for degrees of asymmetry $\lambda = a_{--}/a_{++} = 1$ (the RPM for reference, solid curves), 2, 4, and 6. These values of $\lambda$ correspond to size deviations $|\delta_+| = \delta_- = 0$, $1/2$, $1$, and $2$, respectively. It is evident that ADH theory predicts that size-asymmetry lowers the electrostatic energy, with the effects being greatest at the highest and lowest temperatures. At intermediate $T$ the predictions for varying $\lambda$ are grouped together according to density.

Notable features of the high-$T$ plots of Fig. 2(a) are, first, that the finite values of $u$ attained when $T \to \infty$ are exact. This can be verified by a simple a priori calculation; alternatively, the values may be checked from the exact-order $\rho^0/\sqrt{T}$ term as displayed in Eqs. (2.25) and (2.26) that is correctly reproduced by ADH theory. The limiting slopes at $T^* = \infty$ are discussed further in Ref. [17].

The behavior of $u(T, \rho)$ at low temperatures proves, however, much less satisfactory. From Fig. 2(b) one sees that the energy is predicted to fall increasingly rapidly for increasing asymmetry when $T$ falls, passing well below the ground-state prediction $u = -\frac{1}{2}$ of the symmetric DH theory; but see also [10] for the DHBJDl extensions [9]. Most seriously, however, for large enough asymmetry, the energy drops below the 1:1 size-asymmetric lower bound, namely, $u \approx -1$, established in the Appendix (which also treats general $z_+$ and $z_-$). More concretely, the ground states predicted by ADH theory follow from Eqs. (2.1) and (2.23) which yield

$$u_{ADH}(T=0; \lambda) = -\frac{1}{2}(3 + \lambda). \quad (2.28)$$

This expression evidently violates the bound when $\lambda > 5$ (or $|\delta_+| = \delta_- > \frac{3}{2}$).

Of course, this is unacceptable, but even for smaller values of $\lambda$ this behavior casts doubt on the value of the approximation at low $T$ in the vicinity of the expected critical region. The origin of this behavior can be understood physically by examining the ADH predictions for the mean total charge, say $Q_{z+}^+$, within the subborder zone $a_{++} < r < a_{++}$ that can only be penetrated by like, i.e., positive charges. Recalling the notations in Eq. (2.2) and the approximations (2.4)–(2.8), the theory yields

$$Q_+^z = \int_{a_{++}}^{a_{++}} d^3r \rho_+(r) + \int_{a_{++}}^{a_{++}} d^3r q_+ \rho_+ [1 - \beta q_+ (\phi(r))_+], \quad (2.29)$$

FIG. 2. Effects of size asymmetry on high- and low-temperature energy isochores, according to the linear ADH theory for a 1:1 primitive model. For each density, the RPM isochores (with $\lambda = a_{--}/a_{++} = 1$) are shown as solid curves. The associated, successively lower plots correspond to $\lambda = 2$ (dashed), 4 (dot-dashed), and 6 (dotted, for $\rho^* = 0.1$ only). The density-dependent, $T^* \to \infty$ limiting values are exact. (The near-agreement between two of the curves at $T^* = \infty$ is a coincidence.)

$$B_{2,1} = \pi a_3 z_+ z_- \left[ \frac{\delta_+ (2 + \delta_+ + \delta_- (2 + \delta_-))}{(z_+ - z_-)^2} \right], \quad (2.26)$$

which vanishes for the size-symmetric models (with $\delta_+ = 0$), and

$$B_{2,2} = -\pi a_3 \left[ 1 + \frac{z_+^2 \delta_+ + z_-^2 \delta_-}{(z_+ - z_-)^2} \right]. \quad (2.27)$$
where the precise form of \( \langle \phi(r) \rangle \) follows from the explicit solution of Eq. (2.2) with Eqs. (2.6)–(2.8) [17]. Now, when \( T \to \infty \) the approximation becomes exact, yielding \( Q^\infty_+ = q_+ \rho_+ V^\infty \), where \( V^\infty = \frac{4}{3} \pi (a_+^3 - a_{++}^3) \) is the volume of the subborder zone (see Fig. 1). At finite but large temperatures, Eq. (2.29) correctly predicts the linear decrease of \( Q^\infty_+ \) with \( \beta \) as the (+,+ ) repulsions come into play. Of course, the repulsions become repulsive effective as \( T \) decreases so that, at low \( T \), one expects \( Q^\infty_+ \) to remain positive but to be of magnitude only \( V^\infty q_+ \rho_+ \exp(-\beta a_+^3/Da_{++}) \) (although, because, of the negative screening cloud that builds up for \( r > a_{++} \), this estimate might need to be modified somewhat).

However, if one neglects the screening, the mean potential \( \langle \phi(r) \rangle_+ \) in Eq. (2.29) will be of order \( +q_+ / Da_{++} \), or larger, in the subborder zone. Then, as \( T \) falls and \( \beta \to \infty \) it is evident that Eq. (2.29) is likely to predict, first, a totally unphysical negative value for \( Q^\infty_+ \) (even though no negative charges can enter the subborder zone) and, eventually, a divergence of \( Q^\infty_+ \) to \(-\infty\). Explicit calculations [17] fully bear this out. For example, when \( \lambda = 2 \), and \( \rho^* = 0.01 \), one finds negative \( Q^\infty_+ \) for \( T^* \leq 0.9 \); for \( \rho^* = 0.1 \) \(( \approx \rho_+^* \) \) the change of sign is delayed until \( T^* \approx 0.6 \) but the divergence to \(-\infty \) is more rapid. Since the ADH theory for the border zones embodied in Eqs. (2.5), (2.7), and (2.29) reflects a high-\( T \) expansion, the problems at low \( T \) should not be a great surprise. On the other hand, the successes of the original DH theory for the symmetric case at low \( T \) \(( \text{e.g.,} \ [9,10] \) clearly hinge on the absence of any “charge-unbalanced” border zones.

From which predictions for criticality, etc., follow: see below.

Neglecting all the border zones spares this “original DH” theory the pathology of our ADH theory, but seriously reduces the plausibility of their treatment for size-asymmetric models. Thus, not so surprisingly, the high-\( T / \text{low-} \rho \) expansions of the thermodynamics resulting from Eq. (2.31) differ at the first correction to the limiting behavior—see B.3, in Eq. (2.26)—from the exact results captured by our ADH theory; indeed, the former depends only on the ratio \( x_{-D}^0 \sim \rho / T \), while the exact results are more complex.

### III. MODIFIED ASYMMETRIC DH THEORY

We now discuss a class of modifications of ADH theory (originally introduced in [17]) which avoid the unphysical behavior of the charge density in the border zones: this behavior was identified in Sec. II B as a root cause of the pathological low-\( T \) behavior that included violations of the lower bound on the internal energy (see Appendix). In the following section, we examine the predictions of these modified ADH theories for the critical parameters (including a comparison with the original, 1923 DH proposal for asymmetric systems).

#### A. Introduction of border zone factors

It is clear from the previous considerations of the ADH prediction (2.29) for the charge \( Q^\infty_+ \) in a subborder zone, that the standard DH linearization of the Poisson-Boltzmann exponential is generally inadequate in any zone where the mean charge is necessarily unbalanced because of strongly repulsive short-range, steric repulsions. As an alternative first step, that will still yield an analytically tractable theory, we forego the self-consistent aspect of the Poisson-Boltzmann approximation (2.4) in the border zones and consider replac-
ing the self-consistently determined electrostatic potential \( \langle \phi(r) \rangle_s \) by a fixed but possibly temperature- and density-dependent effective potential \( \phi_\sigma^{\ast}(r,T,T) \).

One possibility is merely to use the bare potential, i.e., to put \( \phi_\sigma^{\ast}=\rho_0^{\ast}/Dr \). The right-hand side of Eq. (2.7) would then read \( q^+_s \rho_+ \exp(-\beta q^+_s/Dr) \); but an obvious drawback of such an approximation is that the resulting closure of Poisson’s equation (2.2) is no longer readily solvable. Instead, we simplify further by neglecting the \( r \)-dependence of \( \phi_\sigma^{\ast} \). Thus we introduce temperature-dependent border zone factors \( F^<_s(T) \) and \( F^>(T) \) and consider the replacement of Eq. (2.7) for a subborder zone by

\[
\langle \rho_q(r) \rangle_+ = \rho^>_q(r,T,T) = q_+ \rho_+ F^<(T), \quad a_+<r<a. \tag{3.1}
\]

Likewise, for a superborder zone (indicated, as before, by the superscript \( \ast \)) we advance

\[
\langle \rho_q(r) \rangle_- = \rho^>_q(r,T,T) = q_+ \rho_+ F^>(T), \quad a<r<a. \tag{3.2}
\]

The calculation of the \( \psi_\sigma \) potentials now proceeds just as in Sec. II but the results take a simpler form, which we distinguish by using a circumflex. Maintaining our convention of a smaller positive ion (\( \delta_+<0 \)) and larger negative ion (\( \delta_->0 \)), we find

\[
\psi^>_+ = \frac{z_+ q_0}{Da} \frac{x_D}{1 + x_D} \left[ 1 + x_D \frac{z_+ T^* \delta_+ F^<_s(T)}{2(z_+ - z_-)} \right]
\times \left[ 2 + \delta_+ - x_D (\delta_+ + \frac{3}{2} \delta^2_+) \right], \tag{3.3}
\]

\[
\psi^>_+ = \frac{z_+ q_0}{Da} \frac{x_0}{1 + x_0} \left[ 1 + x_0 \frac{z_+ T^* \delta_- F^<_s(T)}{2(z_+ - z_-)} \right]
\times \left[ 2 + \delta_- + x_D (\delta_- + \frac{3}{2} \delta^2_-) \right]. \tag{3.4}
\]

These modified potentials, valid for general \( F(T) \), reproduce the symmetric DH theory when \( \delta_\sigma \to 0 \), and always generate the standard limiting laws when \( p \to 0 \). For large \( x_D \propto (p/T)^{1/2} \), these approximations for the \( \psi_\sigma \) behave as \( x_D^2 T^* F \), whereas the standard DH expressions approach constants. While the consequences of this fact are not obvious, some numerical results are discussed in Sec. III C, below.

On the other hand, both of the high-\( T \) second-virial terms \( B_{2,1} \) and \( B_{2,2} \) that are correctly generated by the ADH theory [see Eqs. (2.26) and (2.27)], will be reproduced now if the zone factors satisfy

\[
F^<_s(T) = 1 - \frac{|z_+/z_-|}{T^* (1 + \frac{1}{2} \delta_+)} + O \left( \frac{1}{T^* ^2} \right), \tag{3.5}
\]

\[
F^>(T) = 1 + \frac{1}{T^* (1 + \frac{1}{2} \delta_-)} + O \left( \frac{1}{T^* ^2} \right). \tag{3.6}
\]

In principle, \( F^<_s \) and \( F^> \) could be chosen to generate further exact second-virial terms; but such an approach does not seem of much practical value.

\section{B. Choice of zone factors}

Undoubtedly the simplest reasonable approximation for the zone factors is provided by taking

\[ MF: \quad F^<_s(T) = F^>_s(T) = 1 \tag{3.7} \]

or, equivalently, by dropping the term linear in \( \beta \) in Eqs. (2.5) and (2.7). The results will fail to reproduce the correct \( B_{2,2} \) coefficient in Eq. (2.27), although the exact \( B_{2,1} \) term will be generated. Physically, this approximation amounts to a direct mean-field or van der Waals type approach in which the effects on the internal energy of the \( T=\infty \) unbalanced zone charges, \( Q^<_s = q_+ \rho_+ V^<_s \) and \( Q^>_s = -q_- \rho_- V^>_s \) (where \( V^<_s \) and \( V^>_s \) are the zone volumes), are accounted for in a direct way that neglects fluctuations. The zone charges remain finite and, in fact, fixed, for all \( T \); but, at least for small relative zone volumes \( V^<_s/a^3 = 4 \pi \delta_+ (1 + \delta_+) \) and \( V^>_s/a^3 = 4 \pi \delta_- (1 + \delta_-) \), one might reasonably expect that the initial thermodynamic trends with increasing asymmetry \( \lambda = a_- / a_+ \), for small \( \lambda \), will be correctly predicted.

A second natural step, following our initial discussion, is to take \( \delta^+/r \) to be the direct potential, \( q_0^{\ast}/Dr \), evaluated at the mean radius of the border zones, namely, \( r^= = \frac{1}{2} (a_+ + a_-) \) and \( r^= = (1 + \frac{1}{2} \delta_+) a \). This amounts to adopting the exponential form

\[ \exp^<_s: \quad F^<_s(T) = \exp[-\beta q^+_s/Dr (1 + \frac{1}{2} \delta_+)], \tag{3.8} \]

for the inner zone and similarly for \( F^>(T) \) but with the exponent

\[ y^<_s = + \beta q_+ q_- / Da (1 + \frac{1}{2} \delta_-). \tag{3.9} \]

It is encouraging that these choices precisely satisfy Eqs. (3.5) and (3.6) and so reproduce both \( B_{2,1} \) and \( B_{2,2} \).

As regards the subborder zone where, as discussed in Sec. II B, one expects \( Q^<_s(T,\rho) \) to vanish at low \( T \) (or, at least, approach very small values), the assignment (3.8) seems rather satisfactory. Indeed, combining Eq. (3.1) and the first part of Eq. (2.29) shows that \( Q^<_s \) will remain positive but decrease exponentially rapidly as the \((+,+)\) repulsions “flush out” charge from the subborder zone.

It is worth pointing out that the exponential treatment of the subborder zone can be extended within the DH self-consistent spirit by linearizing the Poisson-Boltzmann factor in Eq. (2.5) about the central value of the direct potential in the subborder zone. Thus, if \( y^<_s \) denotes the exponent in Eq. (3.8), while \( y = \beta q_+ \langle \phi(r) \rangle^<_+ \), one may replace Eq. (2.7) by

\[ \langle \rho_q(r) \rangle^<_+ \simeq q_+ \rho_+ e^{-y} = q_+ \rho_+ e^{-y^<_s} e^{-y} = q_+ \rho_+ e^{-y^<_s} [1 - (y - y^<_s)]. \tag{3.10} \]

The overall factor \( e^{-y^<_s} \) should now ensure the sensible behavior of the subborder zone charge \( Q^<_s \) while the linear factor \( y \simeq \langle \rho_q(r) \rangle^<_+ \) accounts self-consistently for some fluc-
tuation effects while still allowing integration of the closed ADH equations. We have not, however, examined this approach further.

In the superborder zone case, however, matters are significantly different. Certainly, as follows from Eq. (3.6), one expects the mean border-zone charge \( Q^{-}\)(\( T, \rho \)) , which will be positive for \( T=\infty \), to increase initially when \( T \) falls. This is simply because the Boltzmann factor enhances the attractions between the larger, central negative ion and the smaller positive ions. Nevertheless, the growth of \( Q^{-} \) cannot continue indefinitely as would be implied by adopting the \( \text{EXP}^{-} \) form with Eq. (3.9) for \( F^{-} \). Regardless of other effects, the hard cores of the + ions must limit the number that can be packed into a superborder zone at any \( T \); hence \( Q^{-} \) must saturate. This effect may be incorporated into the present framework via a saturation fraction, \( F_{s}^{-}=s \), which prescribes the limiting \( T \to 0 \) superborder zone charge density as \( s q_{+} \rho_{+} \); recall Eq. (3.2). A plausible zone factor is then the “regulated” exponential form

\[
\text{REEXP}^{-} : \quad F_{s}^{-}(T) = \frac{se^{-y^{*}}}{e^{-y^{*}} - 1 + s}, \quad (3.11)
\]

with \( c=s/(s-1) \), which satisfies Eq. (3.6) and approaches \( s \) for large \( y^{*} \) (i.e., low \( T \)). In the limit \( s \to 1 \) (no growth in \( Q^{-} \)) this reduces simply to the mean-field form (3.7).

The difficulty in utilizing this proposal, however, is to know what saturation value \( (s>0) \) is appropriate. At low \( T \), considerations of ion association indicate that the predicted saturation charge in the superzone, namely,

\[
Q_{s}^{-} = V_{s}^{-}q_{+} \rho_{+} s = 4 \sqrt{\pi q_{0}} \delta - \left( 1 + \delta_{-} + \frac{1}{2} \delta_{-}^{2} \right) \frac{1}{|z_{+}/z_{-}|} \rho_{+} s, \quad (3.12)
\]

should not exceed the neutralizing value \( |q_{-}| = |z_{-}|q_{0} \). To meet this condition requires \( s \ll 1/\rho \). While a density-dependent \( s \) and, hence, \( F_{s}^{-} \), can be accommodated without changing Eq. (3.4), the subsequent expressions for the free energy become more complex. Furthermore, when \( \rho_{+} \) increases, \( s \) must decrease and may even fall \( \text{below} \) unity. In these circumstances and in the absence of other effective selection criteria we believe it is appropriate to accept the mean-field form (3.7), i.e., to set \( F_{s}^{-}=1 \) even though \( B_{12} \) will not then be correctly reproduced.

In passing we mention, nonetheless, that we have also explored \( \text{algebraic} \) forms that respect Eq. (3.6) [20]. As an example, one can consider

\[
F_{s}^{-}(T) = (1 + y^{*})\left[ 1 + (y^{*}/s)^{2} \right]^{x}, \quad (3.13)
\]

which saturates at \( F_{s}^{-} = s \) (not necessarily greater than unity) when \( y^{*} = \frac{1}{2} \) but decays to 0 at low \( T \) when \( y^{*} > \frac{1}{2} \). For \( s \geq 2 \) such approximations have an added, unexpected feature, namely, they tend to predict an additional asymmetric critical point at higher densities that does not smoothly connect to the standard DH critical point [9] when \( \lambda \to 0 \). While, owing to the strong short-range interactions implicit in the asymmetric systems, such extra critical points are not obviously unphysical (especially when the diameters are nonadditive), we will not discuss the algebraic forms (3.13) (or \( \text{EXP}^{s} \)) further here.

C. Behavior of energy isochores

Before examining the predictions of the modified ADH theories for criticality, it is instructive to examine energy isochores, such as those plotted in Fig. 3, for several densities near the critical value: compare with the results for the simple ADH theory in Fig. 2(b) but note the difference in vertical scales. At temperatures in the coexistence region \( (T^{*} \leq 0.2) \) and for modest degrees of asymmetry \( (\lambda \approx 3-4) \), both the MF-ADH theory [specified by Eq. (3.7)] and the \( \text{EXP}^{s} \text{MF}^{2} \) ADH theory [where \( F_{s}^{-} \) has been replaced by Eq. (3.8)] predict that the internal energy increases above the RPM value with increasing asymmetry. This reduction in thermodynamic stability suggests, as we will confirm, that the predicted values of \( T_{\text{c}}^{*} \) fall as \( \lambda \) increases.

We also find that the MF-ADH and \( \text{EXP}^{s} \text{MF}^{2} \) ADH theories do, indeed, satisfy the energy bounds of the Appendix for all \( (\rho, T) \) states relevant to the critical and coexistence regions in the additive case. Bound violations can occur, but these arise only at the highest densities \( (\rho_{+} = 1) \) and when the asymmetry is great \( (\lambda > 1) \). While this behavior underlines the two modified ADH theories investigated as overall descriptions of the asymmetric primitive model, the pathologies occur far from the critical region and at unphysically large asymmetry levels.

For moderate to large asymmetry and high density, however, we find that the MF-ADH and \( \text{EXP}^{s} \text{MF}^{2} \) ADH isochores exhibit \textit{nonmonotonic} behavior in \( T \), indicating a thermodynamic instability as previously found in a variety of ion-pairing theories [10]. For both the MF-ADH and \( \text{EXP}^{s} \text{MF}^{2} \) ADH theories, these convexity violations occur...
IV. PREDICTIONS FOR CRITICAL PARAMETERS

The discussions presented above of the various proposed modifications of the asymmetric DH theory indicate that the simpler versions should provide a reasonable basis for further exploration, at least in the case of small size asymmetries. Accordingly, we present here, first, by way of calibration, the predictions of (i) the original, 1923 DH theory embodied in Eq. (2.31); (ii) the MF-ADH theory that uses the simple, mean-field border-zone factors (3.7); and (iii) the EXP\(^\ast\)-MF\(^\ast\) ADH theory that retains the mean-field treatment for the superborder zones but recognizes, via Eq. (3.8), the decrease in subborder zone charge resulting from the Boltzmann-factor-enhanced like-charge repulsions.

The predictions for \(T_c^\ast\) and \(\rho_c^\ast\) for a 1:1 electrolyte are embodied in Fig. 4. The asymmetry variable \(\omega(\lambda) = (1-\lambda)^2/(1+\lambda^2)\) is convenient [14] since it respects the symmetry \(\lambda \leftrightarrow 1/\lambda\). Note that \(\omega(2) = 0.20\), \(\omega(3) = 0.40\), and \(\omega(4.79) = 0.60\), while the point charge limit (\(\lambda = \infty\)) gives \(\omega = 1\).

In Fig. 4 no hard-core terms have been included in the free energy [9]. Thus for the RPM (\(\lambda = 1\), \(\omega = 0\)) one has \(T_c^{\text{DH}} = T_b = 0.0625\) [7,9], which may be compared with simulation estimates yielding \(T_c^\ast = 0.049\) (see [7]). The DH prediction for the critical density is very low, namely, \(\rho_c^{\text{DH}} = 1/64\pi = 0.00497\) [9]; however, this increases to around \(\rho_c^\ast = 0.03\) when Bjerrum ion pairing is introduced [7,9]. Inclusion of hard-core effects, say, via a Carnahan-Starling form, reduces all these parameters by a few percent [9]—and the same is expected to happen for the ADH-based theories.

We note immediately from Fig. 4 that the original DH theory (i) predicts that both \(T_c^\ast\) and \(\rho_c^\ast\) rise rapidly with \(\lambda\). These are precisely the trends found by the MSA (using the energy route) [12] and by the MSA with Bjerrum-Ebeling-Grigo pairing [13]. The modified Poisson-Boltzmann approximations of [11] likewise predict that \(\rho_c^\ast\) increases. (See also Fig. 3 in [14]) In these approximations, however, the initial \(\lambda = 1\) values are well known to be significantly higher \([T_c^\ast(1) = 0.08, \rho_c^\ast(1) = 0.015\text{ to }0.03]\); nevertheless, the proportionate rate of increases are roughly comparable.

By contrast, both of the modified ADH theories predict a strong decrease in \(T_c^\ast\) and \(\rho_c^\ast\) when \(\lambda\) increases from unity: see plots (ii) and (iii). Furthermore, these decreases are in accord with the simulations [14,15] (which, however, start from \(T_c^\ast(1) \approx 0.049\) and \(\rho_c^\ast(1) \approx 0.07\) and the relative rates of fall are quite comparable. For the MF-ADH theory (ii) the critical temperature decreases monotonically and we find \(T_c^\ast(\lambda = \infty) = 0.049\) in the point charge limit \((a_{++} \rightarrow 0)\), while \(\rho_c^\ast\) exhibits a shallow minimum at \(\lambda \approx 4\) and then increases to \(\rho_c^\ast(\lambda = \infty) = 0.006\). By comparison, extrapolation of the simulations (beyond \(\lambda = 6\)) suggests, roughly, \(T_c^\ast(\infty) \approx 0.022\) and \(\rho_c^\ast(\infty) = 0.015\) [14].

In the case of the EXP\(^\ast\)-MF\(^\ast\) ADH theory (iii), however, while \(T_c^\ast(\lambda)\) falls monotonically to about 0.053 when \(\lambda \rightarrow \infty\), the critical density undergoes a shallow minimum around \(\lambda \approx 1.8\) and then rises. Insofar as the simulations seem trustworthy, and exhibit plots that curve downwards (i.e., are concave) vs \(\omega(\lambda)\), it is surprising that the use of the EXP\(^\ast\) choice for \(F^\ast\) leads to apparently inferior predictions. Indeed, on a priori theoretical grounds, the latter would seem superior to the MF assignment \(F^\ast = 1\). We emphasize again, therefore, that various steps in our analysis appear most soundly based when \(\lambda\) is not too large.

FIG. 4. Critical temperature and density predictions for a 1:1 electrolyte with additive hard-sphere interactions as a function of the size asymmetry variable \(\omega(\lambda) = (1-\lambda)^2/(1+\lambda^2)\), which increases monotonically with \(\lambda = a_{--}/a_{++}\). The reduced parameters \(\rho_c^\ast\) and \(T_c^\ast\) are defined via Eqs. (1.4) and (1.5): (i) represents the 1923 DH theory, while modifications of asymmetric DH theory are embodied in (ii) with mean-field factors for both \(F^\ast\) and \(F^\ast\), and (iii) with a Boltzmann-factor (EXP\(^\ast\)) border zone factor for \(F^\ast\) and a mean-field factor \(F^\ast\). The circles denote the \(T_c^\ast\) simulation estimates of Romero-Enrique et al. [14].
For completeness, we report that the MF-ADH and EXP$^\infty$MF$^\infty$ ADH theories predict that the critical ratio $Z_c = p_c/p_{-}\kappa B T$ falls monotonically with $\lambda$ from 0.09036 at $\lambda = 1$ [9], while the original DH theory (Sec. II C) predicts a fall of just a few percent followed, for $\lambda \approx 2$, by a monotonic rise. Inclusion of hard-core terms in the free-energy increases these values (by about 7%) but otherwise the behavior remains similar.

V. SUMMARY AND CONCLUSIONS

We have extended Debye-Hückel (DH) theory to asymmetric two-component hard-sphere electrolytes (i.e., “primitive models”) and computed the predicted critical temperatures and densities. We also have derived, in the Appendix, a lower bound on the internal energy that extends Onsager’s bound [18] and depends only on the valence product $|z_+ z_-|$ and the unlike “collision diameter” $a_{+ -}$. In order to extend the original DH theory [1] (based on the Poisson-Boltzmann equation) to the case of size-asymmetric ions with, say, $a_{- +} > a_{+ +}$, we have identified “border zones” around ions of both species, which prove to be of essential importance. These zones are charge-unequilibrated even at infinite temperature because the larger (negative) ions are geometrically excluded while the smaller (positive) ions may always enter: see Fig. 1.

DH extensions that describe the border zones in a physical way (Sec. III) prove successful in matching trends—as determined by two independent simulation studies [14,15]—in the critical temperature and density with increasing size asymmetry (see Fig. 4 and Sec. IV). This contrasts favorably with other theories, including several based on the mean spherical approximation [11–13], which predict trends opposite to those revealed unequivocally by the simulations.

The existence of the zones complicates the theory in an essential way; however, the usual DH approach can be extended straightforwardly and yields explicit approximations for the internal energy (and, thence, results for other thermodynamic properties). This asymmetric DH (or ADH) theory reproduces the limiting laws and provides the exact high-temperature second-virial coefficients, $B_{2,1}$ and $B_{2,2}$ [see Eqs. (2.25)–(2.27)], down to the point-ion limit.

However, in contrast to the standard DH theory for the symmetric restricted primitive model (RPM) with $\lambda = a_{- -}/a_{+ +} = 1$, the straightforward ADH theory violates the (extended) lower bound on the internal energy in the coexistence region when $\lambda > 5$. Even more seriously, for moderate asymmetries and moderate temperatures, the mean charge in a “subborder” zone (that surrounds $a +$ ion) is predicted to change sign and, at low $T$, to diverge; but, by construction of the model, such behavior is physically impossible. This pathology is readily traced to use of the standard DH linearization of the Boltzmann factor within the border zones: see Sec. II B. Modifications of the ADH theory are thus essential for applications at low $T$.

As shown in Sec. III, one may restore physically sensible behavior while retaining the exact high-$T$ behavior by introducing “border-zone factors,” $F^a(T)$ and $F^b(T)$, originally proposed in [17]. (This also ensures that the energy bounds are no longer violated in the critical region and beyond.) The simplest such modification amounts to a mean-field approach in which the Poisson-Boltzmann factors in the border zones (only) are merely replaced by their $T = \infty$ limits, namely, unity. For a subborder zone (around a smaller + ion) a theoretically preferable approach, dubbed EXP$^\infty$, replaces the self-consistent Poisson-Boltzmann factor by a mean bare-interaction Boltzmann factor [or, better, linearizes about such a mean value: see Eq. (3.10)]. A corresponding exponential treatment of the superborder zone (around a negative ion) proves, however, more problematic owing to physically crucial charge-saturation effects that are hard to elucidate in a precise way. One may expect that both the mean-field and EXP$^\infty$ modifications of linear ADH theory are most reliable at small degrees of asymmetry.

To explore the consequences of the simplest modified mean-field and EXP$^\infty$ ADH theories (just sketched) we have computed the predicted critical parameters as a function of the size-asymmetry $\lambda$ for 1:1 electrolytes [with additive interactions, $a_{+ -} = \frac{1}{2} (a_{+ +} + a_{- -})$]; see Fig. 4. As $\lambda$ increases from unity, the predicted $T_c^a(\lambda)$ and $\rho_c^a(\lambda)$ fall systemati-
cally within both of these modified ADH theories. [See Eqs. (1.4) and (1.5) for definitions of the reduced units.] These decreases accord well with recent simulations [14,15]. On the other hand, an original proposal by Debye and Hückel in 1923, that completely ignores the border zones (see Sec. III C), predicts diametrically opposite trends. Furthermore, current, more sophisticated theories [12,13] make similar predi-
cctions of increasing $T_c^a$ and $\rho_c^a$ (in addition to yielding excessively large values of $T_c^a$ for the RPM [7,9]).

We conclude that DH-based theories seem to extract the basic physics in a quantitatively more reliable way [7,9], even for size-asymmetric systems, than do potentially better, but physically less transparent approaches like the MSA. It is still necessary, however [10], to incorporate Bjerrum ion-pairing and dipole-ion solvation [9] into the modified ADH theories expounded here. (This will also increase the predicted critical densities to better match the anticipated values.) It is not obvious that the correct trends with asymmetry (accepting the validity of the simulations [14,15]) will survive these extensions: it seems likely, nonetheless, that the proper dependence on asymmetry will be preserved (as suggested, e.g., by comparing the results of [12] and [13]).

From a broader perspective, it remains frustrating that more powerful and definitive theoretical techniques have not yet been devised to aid in our understanding of such a fundamental and significant model for condensed matter.

ACKNOWLEDGMENTS

We appreciate the interest, encouragement, and comments of Professor Benjamin P. Vollmayr-Lee, Professor Elliot Lieb, and Professor A. Z. Panagiotopoulos. We are indebted to Dr. S. Banerjee who pointed out an oversight in the Appendix. Professor Katharina Vollmayr-Lee generously translated sections of [1] related to the present work, and Dr. Banerjee and Young C. Kim kindly commented on a draft typescript. The bulk of the research reported here was supported by the National Science Foundation (under Grants...
APPENDIX: ENERGY BOUNDS FOR THE TWO-SPECIES PRIMITIVE MODEL

We here generalize the Onsager bound for the RPM [18] to the general two-component hard-sphere model defined in Sec. II and also develop an (apparently new) lower bound that improves on the simple Onsager result.

The quantity to be bounded is the interaction energy of the ions, $U_{\text{int}}$. Defining the interionic distances $r_{ij}$ and interaction potentials,

$$\varphi_{ij} = q_i q_j / D r_{ij},$$

where $q_i$ is the charge on the $i$th ion, one has

$$U_{\text{int}} = \sum_{i<j} \varphi_{ij} = \sum_{i<j} \sum_{i} u_i^{\text{self}}.$$  \hspace{1cm} (A1)

By adopting the $1/r_{ij}$ form in Eq. (A1), we implicitly assume that the charge densities between any two ions are spherically symmetric and never overlap. To avoid irrelevant singularities, we further assume that the charge density is everywhere finite, i.e., has been suitably “smeared” or distributed.

Following Onsager [18], consider the total electrostatic energy, $U_{\text{tot}}$, which, besides the interaction energy, also includes the self energies, $u_i^{\text{self}}$, i.e., the energies required to assemble the individual ions from an infinitely dispersed charge state of zero energy. As is well known (see, e.g., [21]), the total energy may be expressed in terms of the electric field $E$ as a positive definite quantity, namely,

$$U_{\text{tot}} = \sum_{i<j} \varphi_{ij} + \sum_{i} u_i^{\text{self}} = \frac{1}{8 \pi} \int d^3r |E|^2 > 0. \hspace{1cm} (A3)$$

Note that the smeared charge distributions are assumed to guarantee the finiteness of $U_{\text{int}}$.

1. Generalisation of the Onsager bound

The Onsager bound for the RPM and its direct generalization follow easily from (A3). Rearrangement leads to

$$\sum_{i<j} \varphi_{ij} \geq - \sum_{i} u_i^{\text{self}}. \hspace{1cm} (A4)$$

This inequality holds for arbitrary charge distributions obeying the restrictions stated above; different distributions, of course, lead to different values of $u_i^{\text{self}}$. To obtain the strongest bound, we must minimize the magnitudes of the self energies. This is accomplished by dispersing the ionic charge as much as possible, namely, by placing it in a thin shell on the surface of the largest permissible sphere (of diameter $a_{\text{max}}$). Considering a vanishingly thin shell, the minimal self energy is thus

$$\min (u_i^{\text{self}}) = q_i^2 / 2 D a_{\text{max}}^2. \hspace{1cm} (A5)$$

For the symmetric RPM, the equality of charges, $q_+ = -q_- = q$, and of sizes, $a_{\text{max}}^+ = a_{\text{max}}^- = a$, immediately leads to the original Onsager bound $q^2 / 2 D a$. In the case of nonadditivity, however, one must avoid overlapping charge distributions from distinct ions, so that

$$a_{\text{max}}^i = \min (a_+, a_-). \hspace{1cm} (A6)$$

We now formulate the explicit Onsager bound. First, defining $z = |z_+ / z_-|$, note that the total numbers of particles of each species are

$$N_+ = N / (1 + z) \hspace{1cm} \text{and} \hspace{1cm} N_- = z N / (1 + z).$$

Recalling the definition of reduced energy (1.6) and using the constraint of overall charge neutrality, one finds the bound

$$u(p,T) \geq \mu_{\text{Ons}} = - \frac{z (a a_{\text{max}}^+) + (a a_{\text{max}}^-)}{1 + z} \hspace{1cm} (A7)$$

For the size-symmetric case, where $a_{\text{max}}^+ = a_{\text{max}}^- = a$, the charge asymmetry does not affect the bound in these reduced units, so that $u(p,T) \geq \mu_{\text{Ons}} = -1$. If we separate the three basic size symmetry cases, the result translates into

$$\mu_{\text{Ons}}^I = - \left[ (1 + \delta_+) (1 + z) + 1 / (1 + \delta_-) (1 + z) \right], \hspace{1cm} \text{for} \hspace{1cm} a_+ < a_- < a,$$

$$\mu_{\text{Ons}}^II = - \left[ (1 + \delta_+) (1 + z) + 1 / (1 + \delta_-) (1 + z) \right], \hspace{1cm} \text{for} \hspace{1cm} a_+ < a < a_-,$$

$$\mu_{\text{Ons}}^III = - 1, \hspace{1cm} \text{for} \hspace{1cm} a < a_+ < a_-, \hspace{1cm} (A8)$$

Note that the strongest bound is $-1$, which obtains for Case III, when both the like diameters exceed $a$; this matches the RPM result. On the other hand, $\mu_{\text{Ons}}$ is weaker in Cases I or II, since the bound can then diverge to $-\infty$ when $\delta_+ \to -1$ (or, equivalently, as $a_{\sigma \sigma} = 0$). Physically, shrinking the like diameters (but keeping $a_+$ positive) should not decrease the energy: thus a stronger bound is desirable.

2. Improved bound

To do better we compare the two-species primitive model with another model whose energy can be bounded by an Onsager construction, specifically, an interpenetrating, two-species shell model, consisting of a charge-neutral mixture of uniformly surface-charged spheres of total charges $q_i = z_i q_0$ or $z_i q_0$ and equal diameters $a$, but with no hard-core...
constraint. The shell diameter \(a\) will be identified with \(a_{++}\) for the primitive models. The interaction potential is described further below.

Since the configuration space of the primitive model—in terms of ion-center locations—is a subset of the space of the shell model, any ground-state configuration of the primitive model of energy, say, \(U_{0}^{\text{int}}\), is present in the shell model with energy, say, \(U^{\text{int}}\), which cannot be lower than the shell model ground state, say, \(U_{0.5}^{\text{int}}\). Thus, if we can establish

\[
U_{0}^{\text{int}} = U_{1}^{\text{int}}
\]

(A9)

and also show that \(U_{0.5}^{\text{int}}\) is bounded below, we will obtain a lower bound on \(U^{\text{int}}\), as desired.

We may construct the interaction potential of the interpenetrating shell model, say \(\tilde{\varphi}_{ij}\) (which does not behave as \(1/r_{ij}\) at all separations), as the difference between the total energy of a two-particle system and the sum of the two self energies. Thus if \(\tilde{\rho}_{k}(r, r_{k})\) represents the charge distribution of a shell ion \(k\) centered at \(r_{k}\), and

\[
\tilde{\rho}(r) = \tilde{\rho}_{i}(r, r_{i}) + \tilde{\rho}_{j}(r, r_{j})
\]

(A10)
is the total charge density for two ions, the interaction potential follows from

\[
\varphi_{ij} + \frac{q_{i}^{2}}{Da} + \frac{q_{j}^{2}}{Da} = \frac{1}{8\pi} \int d^{3}r |E|^{2} = \frac{1}{2} \int d^{3}r \tilde{\rho}(r) \phi(r),
\]

(A11)

where \(E(r; r_{i}, r_{j})\) and \(\phi(r; r_{i}, r_{j})\) are the total field and electrostatic potential. If we now define \(\phi_{k}(r; r_{k})\) to be the electrostatic potential resulting from an isolated shell ion \(k\) at \(r_{k}\), we have

\[
\frac{1}{2} \int d^{3}r \tilde{\rho}_{k}(r; r_{k}) \phi_{k}(r; r_{k}) = \frac{q_{k}^{2}}{Da}.
\]

(A12)

Then, using the linearity of the charge density and potential, the relation (A11) may be simplified to yield

\[
\varphi_{ij} = \varphi(r_{ij}) = \frac{1}{2} \int d^{3}r [\tilde{\rho}_{i}(r; r_{i}) \phi_{j}(r; r_{j}) + \tilde{\rho}_{j}(r; r_{j}) \phi_{i}(r; r_{i})].
\]

(A13)

To compare the energy of an arbitrary primitive-ion configuration with that of the corresponding shell-ion configuration in order to establish Eq. (A9), we observe first that because of the pairwise additivity of the interaction energy (A2), one need analyze only two shell ions of the same charge that overlap. To see this, note that all \((+, -)\) ion pairs in a primitive model are separated by distance not less than \(a_{+-} = a\), which is the same diameter as that of the shell ions. Thus, oppositely charged shell ions that correspond to a primitive ion configuration never overlap. The only differences arise when, in the primitive ion system, one or both of the like diameters, \(a_{++}\), are smaller than \(a\). This will allow overlapping shells in the corresponding shell configurations. If \(a_{++}\) and \(a_{--}\) exceed \(a\), the energy of corresponding primitive and shell configurations will always be identical.

Consider then, for concreteness, two positive overlapping shells, separated by a distance \(r_{++}\), with \(a_{++} < r_{++} < a\); see Fig. 5. We want to show that the interaction energy of this pair, \(\varphi(r_{++})\), is less than that for the corresponding nonoverlapping primitive ions which is simply \(q_{+}^{2}/Dr_{++}\).

By symmetry the two terms in Eq. (A13) are now identical. Thus, consider the charge distribution of the right-hand shell in Fig. 5 in the potential of that on the left; the right-hand distribution divides naturally into the two parts shown in the figure: a part exterior to the left-hand shell (bold) and an interior part (dashed). If \(r_{l}\) denotes the position of the left-hand ion, the resulting potential at an exterior point, \(r^{*}\), is simply \(q_{+}/|r_{l} - r^{*}|\) (because these points “see” a spherically symmetric left-hand charge distribution). Conversely, interior points, such as \(r^{*}\), experience only the constant electrostatic potential \(q_{+}/(a/2)\). This is clearly less than the potential they would experience were all the left-hand charge distributed on the smaller primitive ion sphere with the same center \(r_{l}\). Consequently, overlapping like-charged shell ions have a smaller interaction potential than the corresponding primitive ions with the same centers. This establishes the bound (A9).

To obtain a lower bound for the shell model itself, recall Eqs. (A3) and (A10) and bound the total energy of any shell configuration as

\[
U^{\text{int}} = \frac{1}{8\pi} \int d^{3}r |E|^{2} = \frac{1}{2} \int d^{3}r \tilde{\rho}(r) \phi(r) > 0,
\]

(A14)

where the total shell charge density, \(\tilde{\rho}(r)\) can be expressed in the form (A10) but with a sum extending over all the shell ions. The total electrostatic potential, \(\phi(r)\), can be decomposed similarly, yielding

\[
\frac{1}{2} \int d^{3}r \tilde{\rho}(r) \phi(r) = \frac{1}{2} \int d^{3}r \left[ \tilde{\rho}_{1}(r) + \cdots + \tilde{\rho}_{N}(r) \right] \times [\phi_{1}(r) + \cdots + \phi_{N}(r)].
\]

(A15)

Finally, by combining the previously defined shell self-energies (A12) and interaction potentials in Eq. (A13), the inequality (A14) may be rearranged to give
\[ \frac{U^\text{int}}{N} = \frac{1}{N} \sum_{i<j} \Phi_{ij} \geq \frac{U^\text{int}}{N} \geq \frac{1}{N} \left( \frac{N_+ q_+^2}{D_+} + \frac{N_- q_-^2}{D_-} \right) \]

\[ = - \frac{|q_+ q_-|}{D_+}. \quad \text{(A16)} \]

The bound on the primitive model is now completed by combining this with Eq. (A9). In terms of the reduced energy per particle (1.6), the result may be written as

\[ u(r, T) \geq -1. \quad \text{(A17)} \]

Note that the positive definite collision diameter, \( a_+ = a \), and the valencies \( z_\alpha \) do not appear explicitly here since they enter into the definition (1.6). Since \( a_{+-} \) and \( a_{-+} \) are also absent, the bound remains valid for point ions (\( a_{\alpha \sigma} \to 0 \)).