The liquid drop volume, surface and curvature terms and their relationship

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The volume and surface coefficients in the liquid drop expansion share the same physical origin and their physical connection is used to extend the expansion with a curvature term. A possible generalization of the Wigner term is also suggested. This connection between coefficients is used to fit the experimental nuclear masses. The excellent fit obtained with a smaller number of parameters validates the assumed physical connection.

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1. Introduction

In its traditional form, the liquid drop model approximates the binding energy of a given nucleus of mass number \( A \) and charge \( Z \) as [1]:

\[
E_B(A, Z) = -a_v A + a_s A^{2/3} + a_c \frac{Z(Z-1)}{A^{1/3}} + a_a \frac{(A-2Z)^2}{A} \pm \frac{\delta}{\sqrt{A}}. \tag{1.1}
\]

The five terms in this equation are associated with the five independent aspects of nuclei expected to affect the binding energy and are associated with the nuclear volume, surface, Coulomb repulsion, proton-neutron asymmetry, and pairing. The quantities \( a_v, a_s, a_c, a_a, \) and \( \delta \), are the coefficients of the respective terms. A fit of this equation to nuclear masses gives the coefficients and reproduces the experimental values to within 1% or \( \sim 10 \text{ MeV} \) for heavy nuclei.

This result attests to the profound physical content of the overall equation and to the interpretation of its individual terms. The residual 1% discrepancy is due to shell structure. The shell corrections, evaluated according to the Strutinsky procedure [2] and grafted onto the liquid drop model, permit an accurate evaluation of nuclear masses and fission barriers to within 1-2 MeV [3, 4, 5]. This hybrid approach remains to this day the yet unmatched paragon for more sophisticated models such as Hartree-Fock-Bogoliubov [6, 7].

Many additional terms have been suggested, each with their own physical interpretation. An example of this is found in Myers and Swiatecki [3] who suggested:

\[
E_B(A, Z) = -a_v \left(1 - k \frac{I^2}{A^2}\right) A + a_s \left(1 - k \frac{I^2}{A^2}\right) A^{2/3} + a_c \frac{Z(Z-1)}{A^{1/3}} + \frac{W|I|}{A} \pm \frac{\delta}{\sqrt{A}}, \tag{1.2}
\]

where \( I = A - 2Z \). The main difference between equation 1.1 and equation 1.2 is the extension of the neutron-proton asymmetry to the surface energy term. Also, a term linear in \( |I| \) was introduced.

Equation 1.2 implies a connection between volume and surface energies. The authors argued that the change of the volume energy due to the neutron-proton asymmetry \( I \) should be reflected in the surface energy of the system as well, though stating that this was done without empirical evidence [3]. The natural implication is that the surface and volume energies are related to each other through their common origin.

A term linear in \( |I| \) was originally suggested by Wigner in considering the exchange force of nucleons [8]. An empirical observation of such a dependency in the masses was reported by Myers and Swiatecki, hence its addition in the above equation [3].

We extend and generalize the insights discussed above by arguing that the relation between volume energy and the surface energy is strong enough that their coefficients should not be taken as independent variables. Furthermore, we discuss the need of a third term arising from the same physics, proportional to \( A^{1/3} \), in order to create a consistent physical picture of the nuclear binding energy. Finally, a linear term in \( |I| \) is naturally introduced when treating the asymmetry term as the expectation value of the isospin, \( T^2 \).

A revised binding energy equation is fit to the experimental binding energies of the nuclides to test these considerations. The importance of the revised terms is assessed by comparing the fit with the original and revised models.
2. The Liquid Drop Formula: A Truncated Series Expansion

The first term of equation 1.1 called the volume term, and its proportionality to $A$ indicates saturating forces leading to constant density and binding energy per nucleon. The obvious similarity to molecular fluids led naturally to the introduction of the second term, called, just as aptly, the surface term. Its proportionality to $A^{2/3}$ speaks to the lack of saturation on the nuclear surface, whose area, through the constant density of the fluid, should indeed be proportional to $A^{2/3}$. Progressing along the same line, it was widely appreciated that the surface term is a finite size correction and that additional terms in the expansion might be needed, such as a curvature term.

More generally, we can think of a generalized liquid drop formula as a rapidly converging series expansion in powers of $A^{-1/3}$, known as the leptodermous expansion [9]:

$$E_B = -a_v A + a_s A^{2/3} + a_r A^{1/3} + ... \quad (2.1)$$

It is left to be determined how many terms in the expansion are necessary to describe the physics of the nuclear system. The incorporation of a curvature term, with its coefficient $a_r$, proportional to $A^{1/3}$ is almost demanded by the truly small size of nuclei ($A \leq 300$) compared to the size of the drops typically considered in molecular fluids, such as aerosols, where $A \geq 10^6$. Higher order terms also may be of importance due to the small size of nuclei, but would be intractable without an understanding of the lower order curvature term.

The role of the curvature term in nuclear systems was considered only recently. The curvature correction was introduced previously by several authors with ambiguous results [3, 4, 5]. The increased number of parameters and the ability of the traditional liquid drop formula without curvature to fit the data made the problem of identifying the magnitude of this term rather difficult. We believe that it is possible to shed additional light on this subject by considering globally the physical origin of the various terms.

Volume and surface terms both arise from the same physical property of nuclear forces: saturation, and the lack thereof. Thus, surface and volume terms should be related to one another, being themselves different effects of the same cause. Furthermore, the experimental surface and volume coefficients turn out to be approximately equal in magnitude and opposite in sign. Is this an accident or could they possibly be equal?

To answer this question, consider a system of small sticky cubes used to build larger, composite cubes. These cubes interact only when in direct contact. The system is characterized by some bond strength, $\varepsilon$, when two faces are touching. The energy of a cube of $A$ constituents is equal to a volume energy minus a surface energy, just as in the nuclear case. Counting the number of bonds in a cube of size $A$ reveals:

$$E_B^{(cube)}(A) = -3A\varepsilon + 3A^{2/3}\varepsilon. \quad (2.2)$$

Thus, in this model the volume and surface energy coefficients are exactly equal with $a_v = a_s = 3\varepsilon$. This insight motivates setting $a_v = a_s$ without any loss of information.

One difference between this simple model system and a nucleus is the diffuseness of the nuclear surface. What effect does a diffuse surface have on the binding energy of a drop? Since the volume energy is a property of the bulk system it would remain unchanged. The fact that the system naturally becomes diffuse means that it gains a larger binding energy in doing so. The surface
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Table 1: Fits of the nuclear masses with equation 3.3 using different mass ranges and setting $a_v = 0$. All the parameters in units of MeV. The value in the parentheses is the uncertainty in the last digit.

<table>
<thead>
<tr>
<th>Masses</th>
<th>$a_v$</th>
<th>$a_s$</th>
<th>$k$</th>
<th>$a_c$</th>
<th>$\delta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>50-100</td>
<td>15.39(4)</td>
<td>16.81(10)</td>
<td>1.742(7)</td>
<td>0.686(3)</td>
<td>10.3(5)</td>
</tr>
<tr>
<td>100-150</td>
<td>15.39(2)</td>
<td>16.68(7)</td>
<td>1.771(3)</td>
<td>0.6917(14)</td>
<td>12.4(3)</td>
</tr>
<tr>
<td>150-200</td>
<td>15.11(2)</td>
<td>15.66(8)</td>
<td>1.748(3)</td>
<td>0.6760(12)</td>
<td>13.5(3)</td>
</tr>
<tr>
<td>200-250</td>
<td>15.18(6)</td>
<td>15.7(2)</td>
<td>1.768(5)</td>
<td>0.686(3)</td>
<td>13.3(4)</td>
</tr>
</tbody>
</table>

energy would then be lowered in comparison to the sharp surface system. This implies that the surface energy coefficient should be equal to or smaller than the volume energy coefficient, contrary to what is observed in traditional liquid drop fits to the nuclear masses.

As the system is made smaller, more terms in the leptodermous expansion may be needed to properly predict binding energies. If one were to fit the expansion with an insufficient number of terms, what ailments would be observed? The terms included in the equation would have to change from their nominal values to accommodate the lack of higher order terms. Furthermore, the deviation from the nominal value would be worse for smaller masses, where the higher order terms are more important.

Consider fitting the nuclear binding energies to a liquid drop model in various mass ranges. Using a fitting procedure with equation 3.3, which will be described in the following section, table 1 shows the results of such an exercise. Most terms do not vary systematically as the mass range is changed, their apparently random variation being of the order of 1%. The exceptions are the surface energy and the pairing energy. The pairing energy is of unrelated physics and is not discussed here. The surface energy coefficient decreases as the mass range is incremented. This trend indicates that the $A^{2/3}$ term is not sufficient in describing the lack of saturation in the system. As the masses used in the fit increase, the surface term tends to the value of the volume coefficient. Hence, both the need of a curvature term and setting $a_v = a_s$ are motivated.

Now that we see the need for a curvature term, another question arises naturally: What is the origin of the curvature term? In order to answer this question, let us assume a simple liquid with spherical molecules of radius $r_n$. As shown on the left side of figure 1, on a flat liquid surface the molecules should protrude half way on average, thus losing half of their binding energy. If the liquid surface is curved like that of a sphere, as in the right side of figure 1 with a drop radius of $R$, the molecules protrude more, thus losing additional binding. This additional loss increases with increasing curvature. Thus, the curvature and surface terms arise from the same physical effect and their coefficients should be related to one another in a simple way.

In order to obtain a quantitative estimate of this effect, let us consider a model that is essentially geometric in nature. The surface energy is considered to be proportional to the protruding surface area of a constituent residing on the surface times the number of particles present on the surface. As a function of nuclear radius $R$, the resulting exposed surface area, $S$, of a constituent on the surface is:

$$S = 2\pi r_n^2 \left(1 + \frac{r_n}{2R}\right),$$

with the limiting case of a planar system, $S = 2\pi r_n^2$. The number of particles on the nuclear surface
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is proportional to $A^{2/3}$. The overall surface energy is then:

$$ E_s = a_s A^{2/3} \left(1 + \frac{r_n}{2R}\right). \quad (2.4) $$

Since nuclei exhibit a saturation density, the nuclear radius is approximated as $R = r_0 A^{1/3}$, with $r_0$ being a constant. Inserting this relation into equation 2.4 yields:

$$ E_s = a_s A^{2/3} \left(1 + \frac{r_n}{2r_0 A^{1/3}}\right) $$

$$ = a_s A^{2/3} + a_s \frac{r_n A^{1/3}}{2r_0}. \quad (2.5) $$

In this result we identify the usual surface term proportional to $A^{2/3}$ and its surface energy coefficient. Furthermore, we notice a curvature term proportional to $A^{1/3}$ and its curvature coefficient which is dependent on the surface energy coefficient and the ratio of the “molecule” radius to the droplet radius which is directly related to the saturation density.

The above equation is reminiscent of the Tolman correction to the surface tension [10]. This term can be interpreted as the Tolman correction for the nuclear system in its ground state.

Naturally, deviations from sphericity of the molecules would involve a (temperature dependent) reorientation on the surface. This would alter the simple relationship between volume, surface and curvature energies. We will limit the discussion to the case of an isotropic force for the model presented here.

We may check the model further by putting experimental values into equation 2.5. Taking $r_0 \simeq 1.2$ fm [4] and the radius of a free nucleon to be $r_n \simeq 0.9$ fm [11], yields:

$$ a_r \simeq a_s \frac{0.9}{2.4} \simeq \frac{3}{8} a_s. \quad (2.6) $$

It would be notable if a proper fit to nuclear masses were to produce a value close to the above estimate.

To further appreciate the significance of the relation between volume, surface and curvature energies, consider the following. What information is gained in knowing the leptodermous expansion for an arbitrary leptodermous liquid in its ground state?
First, consider the volume energy. The volume energy gives no information of the internal structure of the system. It is just the scale which sets the size of the rest of the terms in the leptodermous expansion.

Now a measurement of the system’s surface energy is made. The particle density of the system can be deduced by comparing the surface and volume energies. This is done by anticipating that the two coefficients will be the same in terms of $A$ and $A^{2/3}$, respectively. Avogadro’s number could thus be inferred.

Finally, the curvature energy is determined and from it the size of a single particle in the liquid can be estimated. This is shown in equation 2.5.

Here we see how the hierarchy of terms in the leptodermous expansion can be related to the internal structure of a fluid. Even though this exercise is pedagogical in nature, it demonstrates the physical significance of each term. It could have allowed Democritus to prove his atomic theory, had he been inclined to do so.

3. Nuclear Mass Fit Results

In the fitting process we use a set of 2076 masses, corrected for shell effects according to Möller et al. [12]. The masses considered in the fits correspond to nuclear masses from reference [13] with $N > 7$, $Z > 7$, and with experimental uncertainties less than 150 keV. The lower limit of neutron and proton numbers is chosen to insure that the included nuclei are large enough to be considered as liquid drops. The restriction on the experimental uncertainties is not only due to the error of the mass, but also to the reliability of the shell correction for masses far away from stability.

The binding energy, $E_B$, of each nucleus is defined as:

$$E_B(A,Z) = Zm_p + (A-Z)m_n - M(A,Z) + \Delta_{\text{shell}}(A,Z),$$

with $m_p$ and $m_n$ being the mass of a proton and neutron, respectively, $M$ is the experimental mass of the nucleus, and $\Delta_{\text{shell}}$ is the shell correction. The liquid drop formula is fit to this binding energy with each nucleus given an equal weight. The mean square deviation of the fit is used to evaluate its goodness:

$$\chi^2 = \sum \left( \frac{E_i^{(ex)} - E_i^{(th)}}{\delta} \right)^2$$

We use the following liquid drop formula:

$$E_B = (-a_vA + a_sA^{2/3} + a_rA^{1/3}) \left( 1 - k \left( \frac{|I|(|I|+2)}{A^2} \right) \right) + a_c \frac{Z(Z-1)}{A^{1/3}} \pm \frac{\delta}{\sqrt{A}},$$

where we insert the mass asymmetry dependence $I = A - 2Z$ both in the volume and surface terms according to Myers and Swiatecki [3]. If the mass asymmetry term is interpreted as an “isospin” dependence, the term linear with $I^2$ should be treated as $T^2$, with $T = |I|/2$. This “isospin” presents itself as the square $T^2$, which we rewrite (with a possibly unjustified quantal sensitivity) as $\langle T^2 \rangle = T(T+1) = |I|(|I|+2)/4$. This has the effect of introducing a linear term in $|I|$ without the addition of a new parameter, as opposed to a freely varying Wigner term [8].

The following fits are performed:
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Table 2: Fits from the four different mass equations as described in the text. All the parameters are in units of MeV. The value in the parentheses is the uncertainty in the last digit.

<table>
<thead>
<tr>
<th>Fit</th>
<th>( a_v )</th>
<th>( a_s )</th>
<th>( a_r )</th>
<th>( k )</th>
<th>( a_c )</th>
<th>( \delta )</th>
<th>( r_n ) (fm)</th>
<th>( \chi^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>15.597(7)</td>
<td>17.32(2)</td>
<td>( a_r = 0 )</td>
<td>1.8048(9)</td>
<td>0.7060(4)</td>
<td>11.4(2)</td>
<td>—</td>
<td>0.58</td>
</tr>
<tr>
<td>B</td>
<td>14.843(3)</td>
<td>( a_v = a_s )</td>
<td>( a_r = 0 )</td>
<td>1.7196(16)</td>
<td>0.6585(4)</td>
<td>10.1(6)</td>
<td>—</td>
<td>4.24</td>
</tr>
<tr>
<td>C</td>
<td>15.25(3)</td>
<td>15.17(17)</td>
<td>3.8(3)</td>
<td>1.779(2)</td>
<td>0.6932(11)</td>
<td>11.3(2)</td>
<td>0.60(5)</td>
<td>0.54</td>
</tr>
<tr>
<td>D</td>
<td>15.264(4)</td>
<td>( a_v = a_s )</td>
<td>3.60(3)</td>
<td>1.7805(8)</td>
<td>0.6938(3)</td>
<td>11.3(2)</td>
<td>0.566(5)</td>
<td>0.54</td>
</tr>
</tbody>
</table>

A. \( a_v \) and \( a_s \) vary independently without a curvature term.

B. Same as above, but forcing \( a_v = a_s \).

C. \( a_v \) and \( a_s \) vary independently with a curvature term.

D. Same as above, but forcing \( a_v = a_s \).

The Coulomb, mass asymmetry and pairing coefficients are left as free parameters in all of the above fits. The results are shown in Table 2 and are discussed below. Figure 2 shows plots of the residual masses of the fits, the exact binding energy with shell corrections included minus the binding energy predicted from the fitted formula.

Comparing fits A and B shows that setting \( a_v = a_s \) without the presence of the curvature term does not ameliorate the situation. Quite to the contrary, the \( \chi^2 \) value is 8 times larger and the plot of the residual masses shows clear deviations. Left without constraint, the surface term incorporates the curvature effects and as a result it becomes larger.

Comparing fits A and C, we observe that the introduction of the curvature term as a free parameter improves the resulting fits as expected. But is the value of \( a_r \) physically meaningful and how does it compare to the expectations of the geometric model?

Rearranging equation 2.6 gives the radius of the nucleon as

\[
    r_n = 2.4 \frac{a_r}{a_s},
\]

in units of fm. Using this equation, the nucleon radius is found to be 0.60(5) fm, smaller than the experimental value of 0.84 fm [11]. This size of deviation is not unexpected from the crude approximations used, and it remains impressive that both the sign and relative magnitude are predicted. Furthermore, the surface energy coefficient moves within error of the volume energy coefficient. The other parameters change within 2% between the two fits, showing consistent results.

By forcing \( a_v = a_s \) with the presence of the curvature correction, as in fit D, the \( \chi^2 \) changes by a fraction of a percent. Also, the parameters not associated with the saturating nuclear force are left unchanged. Thus, no physics is lost with setting \( a_v = a_s \).

The above exercise explains the reason why the volume and surface energy coefficients have previously been treated as independent values. Without taking into account the curvature term, the volume and surface parameters will tend to be irreconcilably different to be considered equal. The addition of the curvature term corrects this discrepancy, and it is found that the surface and volume energies are close to being equal, giving no visible difference in the fitting of the experimental data.
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Figure 2: The residual mass from the corresponding fits. The label in the top left corner of each plot corresponds to the fits listed in the text. The connected lines represent chains of isotopes.

Another fit was performed using $\langle I^2 \rangle = |I|(|I| + x)$, with the added fit parameter $x$. This addition is equivalent to introducing an adjustable Wigner term linear in isospin. Table 3 shows the fit with and without letting $x$ vary. None of the other fit parameters change substantially. As for $x$ itself, it is found to be 1.51(3), which slightly lowers the $\chi^2$ of the fit. With most of the parameters changing less than 1%, the same physics is still captured by setting $\langle I^2 \rangle = |I|(|I| + 2)$.

4. Implications of the curvature term

The presence of a curvature energy, especially important in light nuclei, may imply effects hitherto undiscovered. We give here two examples.

The curvature of the surface in the nuclear deformation landscape, and in particular at the fission saddle point, exhibits large variations going from positive to negative. Therefore, the prediction of fission saddle point configurations and masses will be affected by the presence of a curvature term, which will acquire a tensorial form.
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Table 3: Fits of the nuclear masses to the liquid drop model using different isospin dependencies. The first forces \( \langle I^2 \rangle = |I|(|I| + 2) \), where as the second represents a fit to \( \langle I^2 \rangle = |I|(|I| + x) \). All the parameters are in units of MeV. The value in the parentheses is the uncertainty in the last digit.

<table>
<thead>
<tr>
<th>( a_v )</th>
<th>( a_r )</th>
<th>( k )</th>
<th>( x )</th>
<th>( a_c )</th>
<th>( \delta )</th>
<th>( \chi^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.264(4)</td>
<td>3.60(3)</td>
<td>1.7805(8)</td>
<td>2</td>
<td>0.6938(3)</td>
<td>11.3(2)</td>
<td>0.54</td>
</tr>
<tr>
<td>15.247(4)</td>
<td>3.76(3)</td>
<td>1.7944(10)</td>
<td>1.51(3)</td>
<td>0.6913(3)</td>
<td>11.3(2)</td>
<td>0.46</td>
</tr>
</tbody>
</table>

The fragment distribution predicted by the Fisher model [14] is dependent on the surface energy of the clusters. The theory uses a term proportional to \( A^{\sigma} \) for this purpose. Since the fragment yields are weighted heavily towards lighter fragments away from the critical temperature, the introduction of a curvature term would seem imperative. Furthermore, the curvature term could alter predictions of the critical temperature in a way as yet unknown.

5. Conclusion

Previous efforts have addressed the need of a curvature term in the liquid drop expansion of nuclear masses, but no consistent interpretation was made. Some works state that it is unnecessary, and that it is enough to stop the expansion at the level of a surface term [3]. Other studies give conflicting results, and even the sign of the curvature correction remains ambiguous [4, 5].

We demonstrate that the surface energy coefficient in the traditional liquid drop formula changes when different mass ranges are considered. The decreasing trend in the surface energy coefficient with increasing mass number is consistent with the presence of a curvature term. We present a consistent description of the curvature term’s nature, determine its sign and demonstrate its presence in the nuclear masses.

Simple physical arguments predict that the volume and surface energy coefficients should be equal. Without the introduction of the curvature term, the volume and surface energy coefficients appear to differ from each other. With the addition of the curvature term, the two coefficients agree within error.

The nature of the “Wigner” term linear with isospin is also considered. A slight change in the definition of the squared isospin, possibly quantum mechanical in nature, captures its relative magnitude without introducing an additional parameter.

What is gained through these considerations is a streamlined physical picture of the liquid drop model. Consider the difference of the original liquid drop model in equation 1.1 to the final equation presented here in equation 3.3. Even though the latter appears more complicated, there are the same number of free fit parameters as the former. Instead of adding more and more terms to produce more and more exact representations of the nuclear masses, we have added a geometric physical picture and kept the same number of variables to obtain a more accurate result. The lessons learned with this equation are more telling than letting all the parameters free.

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References