Calculations of Electrostatic Potential in Ionic Crystals with Direct Summation of the Coulomb Potentials of Point Charges

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Electrostatic potentials in ionic crystals have been calculated with direct summation of the Coulomb potential of point charges by the Coker's method. The self-potentials for the lattice ions and the Madelung constants are obtained in high precision and accuracy for several compounds: NaCl, CsCl, CaF₂, ZnS, Cu₂O, 2-4 and 3-3 perovskites. The self-potentials and the Madelung constants for these compounds are consistent with each other and the electrostatic potentials at lattice interstitial points are checked.

KEY WORDS: Electrostatic potential/ Madelung constant/ Self-potential/ Coker's method/ Ionic crystal

I. INTRODUCTION

To understand the chemical bond and the electronic structure of ionic crystals, estimation of electrostatic potentials in the periodic arrangement of the ions is an inevitable task. There are two kinds of methods which carry out the estimation. Direct summation of electrostatic potentials due to the lattice ions is a straightforward method. Several methods have been proposed by Evjen,¹) Frank²) and the others. However, most of the methods are applicable for only special cases. The direct sums are conditionally convergent. For the lattice self-potential for CsCl, for example, the Evjen's method approached to double limits erroneously, as mentioned in the original paper.¹) Moreover, most of the direct-summation methods provide the potentials only at the lattice sites, not at the interstitial sites of the lattice, because these methods depend on symmetries about the sites of consideration. Generally applied methods³,⁴) are on the basis of the indirect method of Ewald.⁵) The ionic charges are distributed in the Gaussian form and the Poisson's equation is solved for the generated potential. Although some parameters are optimized for acceleration of convergence, the indirect methods require great effort and much time of computation.

As the Ewald's method is successful, there must exist an equivalent direct-summation method. A point charge produces the equal potential to that of corresponding ionic charge distribution at positions outside the ion. Hajj has proposed a direct-summation method along with an acceleration procedure of convergence which is applicable to the interstitial sites because of an appropriate symmetric property.⁶) Averaging the potentials which are calculated for some different unit cells are necessary to cancel out the potential due to the surface of the finite

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crystallites. The Hajji’s method is universal but the averaging procedure depends on the crystal structure. Coker has reported a simple and universal method for the electrostatic potentials in the crystals with direct summation.\(^7\) He has discussed that the residual potential due to the surface structure of the finite crystallite causes the trouble in the previous direct-summation methods. He uses asymmetric crystallites where the surface potential at one end cancels out that at the opposite end. The remaining field due to the dipoles lying around the asymmetric point is estimated separately and removed from the lattice sums. Coker has analyzed the convergence polynomials which express the difference between the sums for the finite crystallite and for the infinite crystal. Removal of some lower order terms from the polynomials accelerates the convergence.

The purpose of the present work is to study the Coker’s direct-summation method whose implementation is intended in a molecular orbital computational code. This method is applied to calculations of the self-potentials, the Madelung constants and the electrostatic potential at the interstitial positions of the lattice. The results are examined whether systematic errors occur among the values of several compounds.

II. COMPUTATIONAL METHOD

The lattice self-potential \( \phi_i \) for the \( i \)th ion in the reference cell is

\[
\phi_i = \frac{1}{2} \sum_l \sum_j Z_j e |r_i + r_j - r_l|, \tag{1}
\]

where the Coulomb interactions with the \( j \)th ion owning \( Z_j e \) charges in the \( l \)th unit cell are summed up. The total electrostatic energy \( U \) per mole is expressed as

\[
U = (N_A e^2 / 2x) \sum Z_i \phi_i, \tag{2}
\]

where \( N_A \) is the Avogadro number and \( x \) is the total number of molecules in the unit cell. The Madelung constant \( M_R \) is connected with \( U \) by a characteristic length \( R \) of the crystal structure:

\[
U = -N_A(e^2/R)M_R. \tag{3}
\]

The dimensionless quantity \( M_R \) is calculated with the following equation:

\[
M_R = -(R/2x) \sum Z_i \phi_i. \tag{4}
\]

In the original definition by Madelung, the largest common factor of \( Z_i \)’s is extracted from the Madelung constant in Eq. (4). This factor is included here for the convenience of comparison of the Madelung constant with the self-potential and between the compounds.

The direct-summation method by Coker\(^7\) is briefly summarized. For the unit cell shown in Fig. 1, component ions are a cation and an anion. A space filling procedure using the translational vectors generates crystallites. An asymmetric point with a null electrostatic potential resides at the midpoint between these ions. The pair of ions makes up a dipole and then the method is called the dipolar-addition method. The potential due to the surface structure is cancelled out because of the asymmetric property. The field produced by the dipole \( \mu \) at a distant point \( r \) is expressed as follows:

\[
d\mathbf{E} = \sigma / r^2 \mathbf{d} \mathbf{v}, \quad \tag{5}
\]

\[
\sigma = \mu \mathbf{n} / abc, \quad \tag{6}
\]
where $a$, $b$ and $c$ are the lattice parameters for an orthorhombic crystal. Coker has obtained the solutions of the total field in the crystallites for all the crystal systems.\(^7\) The field for the orthorhombic crystal is estimated as

$$E_x = \frac{8\mu_0}{abc} \tan^{-1} \left( \frac{bc}{a\sqrt{a^2 + b^2 + c^2}} \right).$$

$E_x$ and $E_y$ are obtained from $E_x$ by cyclic permutation. Coker has found that the convergence polynomials for all the crystal symmetries include only even reciprocal powers beginning with the inverse quadratic. Elimination of some lower order terms among successive sums of the crystallites accelerates the convergence.

### III. RESULTS AND DISCUSSION

The self-potentials and the Madelung constants for several compounds are summarized in Table I. In the present calculations, 14 shells around the central unit cell are included. The results contain the decimal figures unchanged between calculations of 14 shells and 15 shells. The first two terms of the convergence polynomial were removed among three successive crystallite sums. For NaCl and CsCl, the Madelung constants in high precision have been reported.\(^6\)\(^7\) The present results agree with them within the significant decimal figures. For simple structures such as NaCl and CsCl, the Madelung constants $M_a$ is equal to the absolute value of the self-potential for the cation and the anion in units of $e/a$, where the lattice parameter $a$ is adopted as the characteristic length for the Madelung constant. These values agree in high precision with each other. No surface potentials contribute to the Madelung constants for these structures because the contribution to the cation site cancels out that to the anion site. The agreement between the Madelung constant and the self-potentials in the present work means that the surface contribution to the potentials at the lattice sites is thoroughly removed.

For ZnS, Cu$_2$O, CaF$_2$, SrTO$_3$ and LaAlO$_3$, the Madelung constants and the self-potentials agree with the reported values and the higher precision is achieved. For comparison of the values among these compounds, alternative estimation was also made. The self-potentials for a compound relate to the Madelung constants for other compounds. For CaF$_2$, these relationships are\(^1\)\(^1\)

$$\phi(Ca) = \frac{M_a(ZnS)}{2},$$

$$= 2M_a(CsCl) + M_a(NaCl).$$

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\[ \phi(F) = \frac{M_a(ZnS)}{2} - M_a(NaCl), \]
\[ = 2M_a(CsCl). \]

For the perovskite structure of \( A^+Z^+(3-z)X_3^{-1} \), they are\(^{12}\)

\[ \phi(A) = Z a_0 - 3a_2 + (3-z)a_3, \]
\[ \phi(B) = (3-z)a_0 - 3a_1 + z a_3, \]
\[ \phi(X) = -a_0 + (3-z)a_1 + (z-2)a_2, \]
\[ a_0 = 3a_1 + 3a_2 + a_3, \]
\[ a_1 = \frac{[M_a(NaCl) - 2M_a(CsCl)]}{6a}, \]
\[ a_2 = \frac{-M_a(NaCl)}{6a}, \]
\[ a_3 = \frac{[M_a(NaCl) + 6M_a(CsCl) - 2M_a(Cu_2O)]}{6a}. \]

These equations relate all the values in Table I. The solutions with the direct summation completely agree with the values derived with the above relations. It is proved that the present values are in high accuracy and precision.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Potential (e/a)</th>
<th>Madelung constant (Ma**)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cation site</td>
<td>Anion site</td>
</tr>
<tr>
<td>NaCl</td>
<td>TW - 3.49512918927</td>
<td>3.49512918927</td>
</tr>
<tr>
<td>CsCl</td>
<td>TW - 2.035361509</td>
<td>2.035361509</td>
</tr>
<tr>
<td>ZnS</td>
<td>TW - 7.565852208</td>
<td>7.565852208</td>
</tr>
<tr>
<td>Cu_2O</td>
<td>TW - 3.782926104</td>
<td>6.476530930</td>
</tr>
<tr>
<td>CaF_2</td>
<td>TW - 7.565852208</td>
<td>4.070723019</td>
</tr>
<tr>
<td>SrTiO_3</td>
<td>TW - 5.387209651</td>
<td>6.455908803</td>
</tr>
<tr>
<td>LaAlO_3</td>
<td>TW - 7.422571160</td>
<td>5.969319576</td>
</tr>
</tbody>
</table>

* Numbers indicate the reference number, TW: this work with the direct summation, TW1: this work using Eq. (8), (10), TW2: this work using Eq. (9), (11), TW3: this work using Eq. (12) - (18).
** The lattice parameter a is the characteristic length for the Madelung constant.
Three kinds of unit cells for CsCl were used to examine the present computational code for the consistency among different crystal systems. The lattice parameters and the atomic positions are as follows.

**Cubic**

\[
a = b = c = 1 \\
Cs: 1/2 \ 1/2 \ 1/2 \\
Cl: 0 \ 0 \ 0
\]

**Tetragonal**

\[
a = b = \sqrt{2} \\
c = 1 \\
Cs: 0 \ 0 \ 1/2 \\
Cl: 1/2 \ 0 \ 0
\]

**Orthorhombic**

\[
a = 2\sqrt{2} \\
b = \sqrt{2} \\
c = 1 \\
Cs: 0 \ 0 \ 1/2 \\
Cl: 1/2 \ 0 \ 0
\]

\[
Cs: 1/4 \ 1/2 \ 1/2 \\
Cl: 0 \ 1/2 \ 0
\]

\[
Cs: 1/2 \ 0 \ 1/2 \\
Cl: 3/4 \ 0 \ 0
\]

\[
Cs: 3/4 \ 1/2 \ 1/2 \\
Cl: 1/2 \ 1/2 \ 0
\]

The self-potentials agree with each other as shown in Table II.

To provide an implement of the present code for molecular orbital calculations, rapid execution is preferable. The self-potentials were obtained with two shells around the central unit cell. The first term of the convergence polynomial expressing the difference between the

<table>
<thead>
<tr>
<th>Crystal system</th>
<th>Potential (e/a)</th>
<th>Cation site</th>
<th>Anion site</th>
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<tr>
<td>Cubic</td>
<td>2.035361509₂</td>
<td>2.035361509₂</td>
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</tr>
<tr>
<td>Tetragonal</td>
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<td>2.035361509₃</td>
<td></td>
</tr>
<tr>
<td>Orthorhombic</td>
<td>2.035361509₄</td>
<td>2.035361509₄</td>
<td></td>
</tr>
</tbody>
</table>

Table III. Comparison in precision of self-potential calculations between crystallites with 14 shells and 2 shells around the central unit cell.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Site</th>
<th>Potential (e/a)</th>
<th>diff. %</th>
</tr>
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<tr>
<td></td>
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<td>14 shells</td>
<td>2 shells</td>
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<td>NaCl</td>
<td>Cl⁻</td>
<td>3.49512918927</td>
<td>3.4955</td>
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<td>CsCl</td>
<td>Cl⁻</td>
<td>2.035361509</td>
<td>2.0350</td>
</tr>
<tr>
<td>ZnS</td>
<td>S²⁻</td>
<td>7.565852208</td>
<td>7.5648</td>
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<tr>
<td>Cu₂O</td>
<td>Cu⁺</td>
<td>-3.782926104</td>
<td>-3.7831</td>
</tr>
<tr>
<td></td>
<td>O⁻²</td>
<td>6.476530930</td>
<td>6.4791</td>
</tr>
<tr>
<td>CaF₂</td>
<td>Ca⁺²</td>
<td>-7.356852092</td>
<td>-7.3576</td>
</tr>
<tr>
<td></td>
<td>F⁻</td>
<td>4.070729019</td>
<td>4.0707</td>
</tr>
<tr>
<td>SrTiO₃</td>
<td>Sr⁺²</td>
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<td>-5.3878</td>
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<tr>
<td></td>
<td>Ti⁺⁴</td>
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<tr>
<td></td>
<td>O⁻²</td>
<td>6.455906803</td>
<td>6.4566</td>
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<tr>
<td>LaAlO₃</td>
<td>La⁺³</td>
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<td>Al⁺³</td>
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<td></td>
<td>O⁻²</td>
<td>5.969919576</td>
<td>5.9698</td>
</tr>
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Table IV. Electrostatic potentials at interstitials in the CsCl crystal.

<table>
<thead>
<tr>
<th>Positions</th>
<th>x/a</th>
<th>y/a</th>
<th>z/a</th>
<th>15 shells</th>
<th>2 shells</th>
<th>Hajj*</th>
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</thead>
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<tr>
<td>0.0</td>
<td>0.25</td>
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<td>0.0</td>
<td>-2.0000000001</td>
<td>-2.00022</td>
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<tr>
<td>0.05</td>
<td>0.25</td>
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<td>-1.9183377003</td>
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<tr>
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<td>0.0</td>
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<tr>
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<td>0.0</td>
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<td>-1.39927</td>
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<tr>
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<tr>
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<td>0.35</td>
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<tr>
<td>0.50</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0000000000</td>
<td>0.000000</td>
<td>0.0000000000</td>
</tr>
</tbody>
</table>

* Ref. 6.

Sums for the finite crystallite and for the infinite crystal was eliminated. The precision is better than 0.04% as shown in Table III. This percentage roughly corresponds to 0.01 eV of the self-potential for real compounds.

For interstitial positions of the lattice, the electrostatic potentials calculated with the present procedure is compared with those obtained with another direct-summation method by Hajj61 (Table IV). Complete agreement is achieved between two methods. The evaluation with two shells around the central unit cell is found enough for the present requirement for the electronic structure calculations. Contours of the electrostatic potential are plotted in the (100) cross section of the CaF2 structure as shown in Fig. 2. This figure has no fault in the symmetrical

![Fig. 2. Electrostatic potential of the (100) plain for CaF2.](image)
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property in contrast to that by the Evjen's method. As an additional remark connecting with the property of this structure, a region of low gradient electric field spreads around the center of the cross section.

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