On Numerical Approximation of Electrostatic Energy in 3D

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Abstract

Approximating the Coulomb self-energy of a charge distribution within a 3-dimensional domain and the mutual Coulomb energy of two charge distributions constitutes often a computational bottleneck in the simulation of physical systems. The present article reports on a recently developed computational technique aimed at the numerical evaluation of the 6-dimensional integrals arising from Coulomb interactions. Techniques from integral geometry are used to show a reduction of the domain from 6-dimensional to 2-dimensional. In the process analytic singularities due to Coulomb’s law are eliminated. Experimental results on the self energy of a charged cube show that the proposed method converges rapidly and is competitive with methods proposed in literature for similar integration problems.
1 Introduction

1.1 The problem

Suppose we are given a domain $D \subseteq \mathbb{R}^3$ in 3-space and a volume charge density function $\rho$ defined in $D$; the electrostatic or Coulomb self-energy of $D$ using the Gaussian unit system is given by the following 6-dimensional integral:

\[ E_D = \frac{1}{2} \int_{p_1, p_2 \in D} \frac{\rho(p_1) \rho(p_2)}{|p_1 - p_2|} \, dp_1 \, dp_2. \]  

(1)

If we are given two domains $D_1$ and $D_2$ in 3-space, endowed respectively with volume charge density functions $\rho_1$ and $\rho_2$, the mutual Coulomb energy is given by the following 6-dimensional integral:

\[ E_{D_1, D_2} = \int_{p_1 \in D_1} \int_{p_2 \in D_2} \frac{\rho_1(p_1) \rho_2(p_2)}{|p_1 - p_2|} \, dp_1 \, dp_2. \]  

(2)

Integrals of this form are found often in physics and chemistry but rarely closed form solutions are known (however a notable exception is reported in the next subsection). Numerical evaluation of integrals (1) and (2) encounters two sources of inefficiency. First of all these integrals are 6-dimensional thus requiring a large number of cubature points. As a rule of thumb, the approximation achieved with $n$ quadrature points in dimension 1, is reached with $n^6$ cubature points in dimension 6, when product quadrature rules are used. Secondly, the integrand function has a singularity whenever the two points $p_1$ and $p_2$ coincide, which happens in integral (1) and may happen in integral (2) as we do not rule out domains that share boundary points. The presence of singularities induces slow convergence in standard numerical integration methods.

In this paper we show that for a vast class of domains and density functions it is possible to transform integrals (1) and (2) so that the kernel is regular and the dimension of the integration domain is reduced to 2, thus making numerical integration an appealing option.

Before we say more about our results we comment on two application areas where such results may be beneficial.
1.2 Applications

1.2.1 Molecular Computations

The well known electron-electron repulsion integral (ERI) is:

\[
(\phi_\mu \phi_\nu | \phi_\lambda \phi_\sigma) = \int_{p_1, p_2 \in \mathbb{R}^3} \frac{\phi_\mu (p_1) \phi_\nu (p_1) \phi_\lambda (p_2) \phi_\sigma (p_2)}{|p_1 - p_2|} dp_1 dp_2,
\]

where \(\phi_\mu, \phi_\nu, \phi_\lambda, \text{ and } \phi_\sigma\) are one-electron orbitals. Such integrals are found in many \textit{ab initio} theories and methods: Hartree-Fock theory and Density-functional theory [1, 2] to mention some of the most important ones. The ERI has the same mathematical structure of the energy integral (2) when we consider as domains \(D_1\) and \(D_2\) the whole space and we interpret \(\phi_\mu \phi_\nu\) (resp. \(\phi_\lambda \phi_\sigma\)) as the function associated with the first (resp. second) domain.

One-electron basis functions are then usually expanded as a linear combination of primitive basis functions. Gaussian type functions [3] have become one of the most popular choices for the basis expansion of atomic orbitals since the pioneering work of Boys [4] showing that the ERI, as well as other relevant integrals, have an analytic exact solution for such class of functions. These analytic solutions are usually obtained through the evaluation of recursive schemes [5, 6, 7], or through the so called Rys polynomial technique [8, 9, 10].

Boerrigter, te Velde and Bearends [11] note that a large number of Gaussian type functions might be needed to tightly approximate one-electron orbitals, thus making the rapid growth of the number of integrals to be evaluated a particularly vexing problem. Other type of basis functions (e.g. Slater Type orbitals, Plane Waves, Bessel functions) may lead to shorter expansion but suffer from the difficulty of analytic or numerical integration. In [11, 12] a cellular approach is used: the space is partitioned in Voronoi polyhedra, where a Voronoi polyhedron is the portion of space closer to a nucleus than any other. Then each polyhedron is split into an inner sphere centered on the atom center and a set of truncated pyramids. Specialized numerical techniques are then used in these two types of domains. Numerical results reported in [11] compare favorably with previously known techniques, notably those based on Diophantine integration [13, 14]. Such a method is suitable for computing particle-distribution interaction integrals, since special attention is paid to singularities at the nuclei, however such technique does not seem to address more general
(6-dimensional) integrals (1) and (2).

A second numerical technique is advocated by Becke [15], (see also [16], [17] and [18]). In [16] integral (3) is split into an external part, corresponding to integration in \( dp_2 \), and an internal part, corresponding to integration in \( dp_1 \) for a fixed \( p_2 \). The internal integral is the potential of the charge distribution \( \phi_\mu \phi_\nu \) at the fixed point \( p_2 \). Such a potential is calculated by considering the equivalent Poisson equation and a finite-difference solution approach. The external integral is then attacked with a technique in [15]. Starting from Voronoi cells based on atomic nuclei Becke defines suitable weighting functions that are continuous, close to the unity within a Voronoi cell and close to zero outside. Using these weight functions, an arbitrary 3-dimensional integral can be reduced to a sum of atom-centered integrals, for which product quadrature rules in spherical coordinates are used.

Further refinements and tuning of the approaches in [11] and [15] for three-dimensional integrals are investigated in [19]. The approach in [20, 21] to the evaluation of integrals of potential theory has some high level similarity with that of Becke, although in a different context.

1.2.2 Energy calculations for crystals

In several models of matter we can distinguish a discrete component made of charged point particles and a continuous component made of continuous distributions of charge (see e.g. [22]). Thus formally we can split the total electrostatic energy into the contribution of the point charges, the mutual energy due to the interaction of point charges and the distribution of charge, and finally the contribution of the distribution of charge. The first contribution is expressed formally as a double summation of the Coulomb energy over pairs of particles. Several techniques, ranging from fast \( n \)-body methods to periodic boundary conditions (Madelung sums [23], Ewald summation [24]), are available to speed up the computation. The second contribution involves a sum of 3-dimensional integrals. Numerical techniques for such integrals have been mentioned above in the context of DFT calculations. In the special case of uniform distribution in a cube some analytic solutions are also known [25, 26]. For the energy of distribution of charges, which are represented by 6-dimensional integrals of type (1) or (2), save the above mentioned references, there is a notable lack of specific
techniques available.

In the study of crystal or quasicrystal lattices it is customary to associate each particle with a convex polyhedron containing the particle. Besides Voronoi cells, space filling polyhedra are also used [27, 28]. Moreover we may want to associate a charge distribution to each polyhedron (e.g. to maintain electroneutrality). Although a constant distribution of charge is a reasonable first choice, more precise models might include non-constant distributions to fit known data (either experimental or obtained through auxiliary computations). Thus energy calculations for regular crystal lattices (not necessarily with cubic symmetry), or even irregular lattices do conceivably benefit from general techniques for computing 6-dimensional energy integrals over convex polyhedral domains with non-constant densities of charge.

1.3 Main results, experiments and comparisons

The main contribution of this paper is a general efficient method to compute integrals (1) and (2) when:

- The domain \(D\) (resp. \(D_1\) and \(D_2\)) is a compact convex polyhedron.
- The density \(\rho\) (resp. \(\rho_1\) and \(\rho_2\)) is a polynomial function in Cartesian coordinates.

As we noted above convex polyhedral domains arise naturally in methods based on Voronoi tessellations or on space-filling polyhedra. Although most popular basis functions are not polynomial (usually exponential terms are present), piecewise polynomial functions may be used to fit any given function in 3-dimensional space.

The proposed method has been implemented and tested on approximating the self-energy of a uniformly charged cube (a more detailed description is in Section 5). A reference value is obtained by a high-order Gaussian integration of the formula in [26] for the potential of a cube at a point. As a focus for comparison we concentrate on the following experimental result: using 1,000 Gaussian points our method attains an absolute error between \(10^{-5}\) and \(10^{-6}\), without exploiting the symmetries of the cube. If we exploit explicitly symmetries of the cube, for the same number of points, the absolute error of our method is in the range between \(10^{-9}\) and \(10^{-10}\).
In [11, p. 103] it is reported an accuracy of $10^{-3}$ using a number of points per polyhedron in the range from 700 (for hydrogen) to 3,000 (for uranium), and exploiting symmetries. However, the integrals considered in [11] are 3-dimensional while ours are 6-dimensional before the geometric transformation. Results in [12, pp. 95-96] on computing the overlap integrals (3-dimensional) for a Slater type function in a convex polyhedron show an error in the range $[10^{-6}, 10^{-7}]$ for a number of points from 2,500 points for small molecules with many symmetries up to 264,000 for large molecules without symmetries.

In [15] it is reported a precision in the range from $10^{-4}$ to $10^{-5}$ using a grid of integration points of size equal to or greater than $20 \times 50 \times 50 = 50,000$ for the integration of three-dimensional functions whose analytic closed form is known.

Our preliminary experiments and comparisons with results in [11, 12, 15] indicate that our method for six dimensional integrals can attain performances comparable to those of competing methods even when applied just to 3-dimensional integrals.

1.4 Integral Geometry and Computational Geometry

Our result is obtained by applying to integrals (1) and (2) geometric transformations already applied successfully to other problems ranging from radiosity (approximation of form factors [29]), to calculation of electrostatic forces [30] and the boundary element method (entries of the stiffness matrix for systems of conducting bodies [31, 32]).

The first step of the transformation involves transforming integrals (1) and (2) into integrals over lines in 3-space. The second step consists in choosing a particular form of the differential measure of lines in 3-space so that we can separate our integral into an external integral over the set of directions, and an internal integral which can be evaluated analytically. The new kernel is evaluated using methods from computational geometry. As a result the initial 6-dimensional integral is reduced to a two-dimensional one. Moreover, while the original kernel in (1), (2) and (3) is singular, the new kernel is regular everywhere in the domain of integration. Although the general scheme is the same as in the above mentioned results, the transformation depends critically on the exponent of the factor $|p_1 - p_2|$ thus requiring in this paper a new derivation starting from first principles.

\footnote{Such integrals are expressed in spherical coordinates, thus with a linear and two angular parameters.}
1.5 Organization of the paper

The paper is organized as follows. In Section 2 we give the geometric transformation of integrals (1) and (2) for any convex compact polyhedron endowed with a uniform charge density. In Section 3 the result is extended to any distribution of charge polynomial in Cartesian coordinates. In Section 4 we describe the overall algorithm. In Section 5 we discuss the implementation, experiments, and numerical results.

2 Geometric Transformation: uniform charge density

In this section we assume that the domain $D$ of interest is a convex and compact body $B$ in three-dimensional space, endowed with a uniform charge density $\rho$. We assume the space to be vacuous or filled with a homogeneous non-polarizable medium, and we assume a fixed coordinate system. The purpose of this section is to use tools from integral geometry and differential calculus in order to rewrite equations (1) and (2) in a form convenient for numerical integration. We begin by considering the electrostatic potential generated by the body $B$ at a point $p$ of the space:

$$V_B(p) = \rho \int_{q \in B} \frac{1}{|p-q|} dq.$$  

The idea is to express $V_B(p)$ as a weighted integral over the set of straight lines passing through $p$. The weight of each line $L$ is given by the charges lying on $L$, and it can be expressed in terms of the length of the intersection of $L$ with the body $B$.

2.1 Preliminaries

Let us introduce some notation, as well as recall some elements of differential calculus and integral geometry (see e.g. [33] and [34]). The first step is to introduce the set of straight lines in space and define a measure on it. We denote with $\mathcal{L}$ the set of straight lines in three dimensional space. Given a point $p$, $\mathcal{L}_p$ is the set of lines $L \in \mathcal{L}$ passing through $p$. We will use $L$ for lines in either $\mathcal{L}$ or $\mathcal{L}_p$, but to make the expressions clearer we will denote their differential measures of lines respectively with $dL$ and $dL_p$. A straight line can be identified in a number of different ways, depending on the coordinates used. To determine a line in
$L \in \mathcal{L}_p$ we only need to specify a direction $u$, so that $L \equiv L(u)$. For a generic line $L \in \mathcal{L}$, for reasons that will become apparent soon, we specify a direction $u$ and the intersection $s$ between $L$ and a plane $S_u$ orthogonal to $u$, so that $L \equiv L(s, u)$. Note that this is just one particular parameterization of lines in the space.

Santaló [34] explains how to find a density for subsets of $\mathcal{L}$ which is invariant under the group of rigid motions; moreover, this density is unique up to a constant factor. The density for straight lines in the space is simply

$$dL = ds du,$$

where $du$ is the differential measure of directions and $ds$ is the surface element on a plane $S_u$ normal to $u$. This nice relation justifies our choice of $(s, u)$ to determine $L$. Other representation of lines produce more complicated expressions.

The direction $u$ corresponds to a point on the surface of a unit sphere. However it will be convenient to identify antipodal points, so that we will really be working on a hemisphere denoted with $\frac{1}{2}\Omega$ whose measure is thus $2\pi$. If we express the direction $u$ in polar coordinates $(\theta, \phi)$ then it holds $du = \sin\theta d\phi d\theta$. For the set $\mathcal{L}_p$ we have simply $dL_p = du$. Intuitively, this differential element can be seen as a small cone with vertex $p$, extending in both directions from $p$, where $du$ equals the solid angle at $p$. The measure of the entire set $\mathcal{L}_p$ is $\int dL_p = |\frac{1}{2}\Omega| = 2\pi$. The physical dimension of $dL$ is [length$^2$], while $dL_p$ is adimensional.

Now we relate these differential elements to the element of volume $dq = dx dy dz$ at the point $q$ of a body $B$. In fact this will enable us to rewrite in a more geometric fashion the classical potential formula. Fix a point $p$, and let $r = |p - q|$ be the distance between $q$ and $p$; then it holds that

$$dq = r^2 dr dL_p. \tag{5}$$

This can be obtained by using polar coordinates, writing $x = r \sin\theta \cos\phi$, $y = r \sin\theta \sin\phi$, $z = r \cos\phi$, and applying the rules of exterior calculus. It is also easily seen geometrically because $dq$ is approximated by a cylinder with base $r^2 dL_p$ and height $dr$ [33].

The other important relation that we need relates the (exterior) product of two differential volumes to the differential of lines. Let $L$ be the line passing through points $p$ and
and let \( r_1 \) (resp. \( r_2 \)) be the distance of \( p \) (resp. \( q \)) from a fixed point of reference on \( L \).

The following holds [34, p. 237]:

\[
dp dq = |r_1 - r_2|^2 \, dL \, dr_1 \, dr_2.
\] (6)

### 2.2 Electrostatic potential at a point

We are now ready to find a formula alternative to (4) for the potential field generated by a convex body \( B \) with uniform charge density. The intuition behind the following theorem is that we take a differential cone with vertex \( p \) and sum up the contributions of all the charges lying in the cone. We find their total contribution to be \( (r_{\text{max}}^2 - r_{\text{min}}^2) dL_p \). Next we integrate over all the directions \( L_p \). The effect due to a fixed differential charge is counted exactly once, because there is only one line \( L \in \mathcal{L}_p \) passing through it.

**Theorem 1** For a line \( L \in \mathcal{L}_p \), let \( \ell = |L \cap B| \) and \( m = |L \cap C| \), where \( C \) is the convex hull of \( B \) and \( p \). Then the potential \( V_B(p) \) generated by \( B \) at a point \( p \) external to \( B \) is

\[
V_B(p) = \frac{\rho}{2} \int_{L \cap B \neq \emptyset} \ell (2m - \ell) \, dL_p,
\] (7)

where the integral is over the set \( \mathcal{L}_p \). If \( p \) is inside \( B \), the potential at \( p \) is

\[
V_B(p) = \frac{\rho}{2} \int_{\mathcal{L}_p} (\ell_1^2 + \ell_2^2) \, dL_p,
\] (8)

where \( \ell_1 \) and \( \ell_2 \) are the lengths of the two segments in which \( p \) splits \( L \cap B \).

**Proof.** Let \( p \) be external to \( B \), and fix a line \( L \) through \( p \) and intersecting \( B \). Clearly \( p \notin L \cap B \). We denote with \( r_{\text{min}} \) and \( r_{\text{max}} \) the minimum and maximum distance of \( p \) from the points of \( L \cap B \). Now the potential at \( p \) is given by the classical formula (4), which in view of (5) becomes

\[
V_B(p) = \int_{q \in B} \frac{\rho}{|p - q|} \, dq = \rho \int_{L \in \mathcal{L}_p} \int_{q \in L \cap B} r \, dr \, dL_p,
\]

where \( r = |p - q| \). For a fixed line \( L \) intersecting \( B \), we have \( \int r \, dr = (r_{\text{max}}^2 - r_{\text{min}}^2)/2 \) because \( r \) varies between values \( r_{\text{min}} \) and \( r_{\text{max}} \). If \( L \cap B = \emptyset \) then \( \int r \, dr = 0 \). Consequently we can integrate on the domain \( \{ L \in \mathcal{L}_p : L \cap B \neq \emptyset \} \), thus obtaining:

\[
V_B(p) = \frac{\rho}{2} \int_{L \cap B \neq \emptyset} (r_{\text{max}}^2 - r_{\text{min}}^2) \, dL_p.
\]
Now the first part of the theorem follows by substituting \( \ell = r_{\max} - r_{\min} \) and \( m = r_{\max} \) and rearranging the formula. This concludes the proof of the first part of the theorem.

When \( p \) is inside the body \( B \), the previous reasoning applies to both directions of a single line \( L \), where \( r_{\min} = 0 \) and \( r_{\max} = l_1, l_2 \) respectively. Moreover, all lines in \( \mathcal{L}_p \) intersect \( B \). This proves the second part of the theorem. \( \square \)

Formulas (7) and (8) have the important property of having a regular kernel, while in the classical formulation (4) the kernel may diverge.

2.3 Self-energy of a body and mutual energy of two bodies

Now we apply the result above in order to find the potential energy of a body or system of bodies. Intuitively, we will find that for a fixed differential element \( dL \) (which can be imagined as a “fat cone”) the contribution to the total potential energy given by the interaction between charges in \( B \cap dL \) is \( \rho \ell^3 dL/6 \). We will obtain the total energy by integrating all the differential contributions. The contribution of two fixed particular differential charges is counted exactly once, since two points define a unique straight line in the space.

**Theorem 2** The potential self-energy \( E_B \) of a convex body \( B \) with uniform charge density \( \rho \) is given by

\[
E_B = \frac{\rho^2}{6} \int_{L \cap B \neq \emptyset} \ell^3 \, dL, \tag{9}
\]

where \( L \in \mathcal{L} \) is a straight line in the space and \( \ell \) is the length of the intersection \( L \cap B \).

**Proof.** Let us start from the classical expression (1) for the potential energy of a body \( B \), where \( \rho \) is constant and we use relation (6):

\[
E_B = \frac{\rho^2}{2} \int_{p,q \in B} \frac{1}{|p - q|} \, dp \, dq
= \frac{\rho^2}{2} \int_{p,q \in B} |p - q| \, dL \, dr_1 \, dr_2,
\]

where \( L \) is the line passing through \( p \) and \( q \), and \( r_1, r_2 \) are the distances of \( p, q \) from a fixed point on \( L \). We can exchange the order of integration, integrating first over the lines \( L \) intersecting \( B \) and then over the pair of points laying on \( L \). We obtain

\[
E_B = \frac{\rho^2}{2} \int_{L \cap B \neq \emptyset} \left( \int_{p,q \in B \cap L} |p - q| \, dp \, dq \right) \, dL.
\]
It can be easily seen that the inner integral evaluates to $\ell^3/3$, and the theorem follows. □

The same result can be obtained starting from the well-known relation $E_B = (1/2) \int \rho(p) V_B(p) \, dp$ and applying formula (8) for the potential at a point.

The value $I_3 = \int 3 \, dL$ is a geometric invariant of the object $B$, and the theorem we just proved is the three-dimensional case of a general relation which can be found in [34]. Using a formula from integral geometry [34, p. 231] we can obtain the bound $I_3 \leq (3/2) \mathcal{V}^2 \mathcal{A}$ (where $\mathcal{V}$ and $\mathcal{A}$ denote respectively the volume and the superficial area), and this translates directly into a useful bound for $E_B$.

**Theorem 3** The potential energy $E_{B_1, B_2}$ of two convex bodies $B_1, B_2$, each with uniform charge density $\rho_1, \rho_2$, is given by

$$E_{B_1, B_2} = \frac{\rho_1 \rho_2}{2} \int_{L \in \mathcal{L}_{12}} \ell_{B_1} \ell_{B_2} (2t - \ell_{B_1} - \ell_{B_2}) \, dL,$$

where $\ell_{B_1} = |L \cap B_1|$, $\ell_{B_2} = |L \cap B_2|$, $t = |L \cap C|$, $C$ is the convex hull of $B_1$ and $B_2$, and $\mathcal{L}_{12}$ is the set of lines which intersect both $B_1$ and $B_2$.

**Proof.** We consider expression (2) and proceed like in the previous theorem. We obtain

$$E_{B_1, B_2} = \rho_1 \rho_2 \int_{\mathcal{L}_{12}} \left( \int_{p \in B_1 \cap L, q \in B_2 \cap L} |p - q| \, dp \, dq \right) \, dL.$$  

By means of some calculus we obtain that the inner integral evaluates to $(1/2) \ell_{B_1} \ell_{B_2} (2t - \ell_{B_1} - \ell_{B_2})$ and the theorem follows. □

### 3 Geometric Transformation: non uniform distribution of charges

In this subsection we will extend the theory presented in Section 2 to the case of arbitrary distribution of charges. We will obtain formulas involving integrals over the set of straight lines, where each line is assigned a “weight” which depends on the body under study.

Consider first the electrostatic potential at a point $p$. For a generic distribution of charge on $B$ formula (4) becomes:

$$V_B(p) = \int_{q \in B} \frac{\rho(q)}{|p - q|} \, dq.$$
Proceeding like in the proof of Theorem 1 we can rewrite the last expression as

\[
V_B(p) = \int_{L \cap B \neq \emptyset} \int_{q \in L \cap B} \rho(q) r \, dr \, dL_p
= \int_{L \cap B \neq \emptyset} w_{B,p}(L) dL_p,
\]

where we defined \( w_{B,p}(L) = \int_{q \in L \cap B} \rho(q) r \, dr \). The quantity \( w_{B,p}(L) \) can be thought of as the “weight” of line \( L \), and represents its contribution to the potential at \( p \) due to \( B \). We already saw that \( w_{B,p}(L) \) can be computed explicitly if \( \rho(q) \) is constant. In the general case, we apply a change of coordinates as follows. Consider an orthogonal system of coordinates \((x', y', z')\) which has the origin at \( p \) and the axis \( z' \) parallel to \( L \). We rewrite the density function in this system of coordinates \( \rho(p) = \sigma(x', y', z') \). Notice that the coordinates \( x' \) and \( y' \) are the same for all the points on \( L \), and we will write simply \( \sigma(z') \equiv \sigma(x', y', z') \).

The weight of line \( L \) is thus

\[
w_{B,p}(L) = \int_{q \in L \cap B} \sigma(r) r \, dr.
\]

Denoting with \( \tilde{\sigma}(x) \) an antiderivative of \( \sigma(x) \), and with \( \tilde{\sigma}(x) \) an antiderivative of \( \tilde{\sigma}(x) \), it is easily seen that \( F(x) = \tilde{\sigma}(x)x - \tilde{\sigma}(x) \) is an antiderivative of \( \sigma(x)x \). Finally, let \( \ell_1 \) and \( \ell_2 \geq \ell_1 \) be the \( z' \) coordinates of the extreme points of \( L \cap \partial B \). We obtain for \( w_{B,p}(L) \) the expressions

\[
w_{B,p}(L) = \begin{cases} 
F(\ell_2) - F(\ell_1), & \text{if } p \notin B; \\
F(\ell_1) + F(\ell_2) - 2F(0), & \text{if } p \in B.
\end{cases}
\]

Consider now the self-energy of a body \( B \), given by expression (1). In the same manner as before we obtain

\[
E_B = \int_{L \cap B \neq \emptyset} \int_{p,q \in L \cap B} \frac{1}{2} \rho(p)\rho(q)|p - q| dpdq dL
= \int_{L \cap B \neq \emptyset} w_B(L) dL,
\]

where we defined

\[
w_B(L) = \int_{p,q \in L \cap B} \frac{1}{2} \rho(p)\rho(q)|p - q| dpdq.
\]

We can apply the same linear transformation as above to express \( \rho(p) \) in a coordinate system \((x', y', z')\) where \( z' \) is parallel to \( L \) (the origin can be fixed arbitrarily). We denote
the transformed function with $\sigma(x', y', z) \equiv \sigma(z')$, and define $\tilde{\sigma}(x)$ and $\tilde{\sigma}(x)$ as before. Let $\ell_1$ and $\ell_2 \geq \ell_1$ be the $z'$ coordinates of the extreme points of $L \cap \partial B$. Then the weight of $L$ can be written as

$$
\begin{align*}
\omega_B(L) &= \int_{r_1, r_2 = \ell_1}^{\ell_2} \frac{1}{2} \sigma(r_1) \sigma(r_2) |r_1 - r_2| dr_1 dr_2 \\
&= \int_{r_1 = \ell_1}^{\ell_2} \sigma(r_1) [\tilde{\sigma}(r_1) - \tilde{\sigma}(\ell_1)(r_1 - \ell_1) - \tilde{\sigma}(\ell_1)] dr_1 \\
&= \tilde{\sigma}(\ell_2) \tilde{\sigma}(\ell_2) + \tilde{\sigma}(\ell_1) \tilde{\sigma}(\ell_2) - \tilde{\sigma}(\ell_1) \tilde{\sigma}(\ell_2) - \tilde{\sigma}(\ell_1) \tilde{\sigma}(\ell_1) + \\
&- \tilde{\sigma}(\ell_1) \tilde{\sigma}(\ell_2)(\ell_2 - \ell_1) - \int_{\ell_1}^{\ell_2} \tilde{\sigma}(x)^2 dx
\end{align*}
$$

(11)

It is an easy observation that for constant $\rho$ this formula gives $\omega_B(L) = \rho^2 (\ell_2 - \ell_1)^3/6$, as we obtained in Theorem 2.

The case of two interacting bodies is in all similar to the one-body case, the only difference being in the integration limits for $r_1$ in expression (11).

4 Algorithm for the evaluation of the integrals in polyhedral domains

The formulas obtained so far for electrostatic quantities are not in a computable form because of the presence of the differential $dL$. In this subsection we will choose a particular parameterization of lines that is suitable for computation. We will obtain two-dimensional integrals with a smooth kernel. Moreover, we will explain how to compute this kernel exactly by analytic integration when the bodies are polyhedral objects and the charge distribution is polynomial in $(x, y, z)$. To keep the discussion clear, we will focus our attention on the computation of $E_B$, given by formula (9) or its generalization (10).

As we explained in subsection 2.1, we choose to represent a line $L$ in 3-space as a pair $(s, u)$, where $u \in \frac{1}{2}\Omega$ and $s \in S_u$. Denote with $B_u$ the projection of the body $B$ onto the plane $S_u$. A line $L \equiv (s, u)$ intersects $B$ iff $s \in B_u$. So the energy of $B$ in expression (10) can be written

$$
E_B = \int_{u \in \frac{1}{2}\Omega} \int_{s \in B_u} \omega_B(s, u) ds du = \int_{\frac{1}{2}\Omega} K(u) du,
$$

(13)
where
\[ K(u) = \int_{s \in B_u} w_B(s, u) ds. \]  

We integrate numerically over the set \( \frac{1}{2} \Omega \) by approximating the integral with a weighted sum of values of the kernel \( K(u) \) at selected points in the integration domain. The domain \( \frac{1}{2} \Omega \) is a suitable domain of integration because a number of results exist on generating distributions of points on the sphere [35]. In particular we can map the sphere into a rectangular domain using spherical coordinates \((\theta, \phi)\); however to simplify certain formulas we choose coordinates \((z, \phi)\), where \( z = \cos \theta \), so that the differential element of directions becomes \( du = \sin \theta d\theta d\phi = dz d\phi \).

Next we show that the value of \( K(u) \) for any fixed given value of \( u \) can be computed exactly via analytic (non numerical) integration.

We compute the value of \( K(u) \) for polyhedral domains and polynomial distributions by means of the following algorithm. The key observation is that the values \( \ell_1 \) and \( \ell_2 \) in the expression for \( w_B(L) \) are piecewise linear, and that \( w_B(L) \) is a piecewise-polynomial function.

1. Fix an orthogonal coordinate system \((x', y', z')\), where the \( z' \) axis is parallel to the direction \( u \), and \((x', y')\) span the plane \( S_u \); next, orthogonally project the edges of the polyhedron \( B \) over \( S_u \).

2. Compute the transformed \( \sigma(x', y', z') \equiv \rho(x, y, z) \) by means of a change of variables; if \( \rho \) is a polynomial in \((x, y, z)\), then also \( \sigma \) is a polynomial in \((x', y', z')\); thus, it is possible to compute symbolically \( \bar{\sigma}, \bar{x} \) and \( f \bar{\sigma}^2 \).

3. Compute the partition of the plane induced by the projected edges; to this purpose a variety of algorithms exist in computational geometry literature [36, 37]. The work required at this step is \( O((n + k) \log n) \), where \( n \) is the number of edges and \( k \) the number of intersections between projected edges, using a method of Bentley and Ottman [38];

4. The global integral \( K(u) \) can be obtained by summing the quantities \( K_f(u) = \int_{s \in f} w_B(s, u) ds \), where \( f \) ranges among all the faces of the planar decomposition;
5. For each face $f$ in the planar decomposition, the quantities $\ell_1$ and $\ell_2$ in (12) are linear functions of $x', y'$; so their analytic expression can be interpolated from their values at the vertices of $f$;

6. Since $\ell_1$, $\ell_2$ and $\sigma$ polynomials, the function $w_B(s, u)$ is also a polynomial, whose expression can be computed analytically;

7. Apply Green's theorem to compute the value $K_f(u)$ using only the values of $w_B$ at the vertices of face $f$;

8. Finally, compute $K(u) = \sum_f K_f(u)$.

The values of $\ell_1$, $\ell_2$ at the vertices of the planar decomposition can be computed by visiting the graph representing the decomposition. The procedure described above applies when $B$ is a polyhedron and $\rho$ is a polynomial, however it applies also to any another function for which easy algebraic manipulation is possible. Notice that these two conditions influence two different aspects of the computation: if $B$ is not a polyhedron, but we can compute $\ell_1$, $\ell_2$ exactly, we can still compute $w_B(L)$ and perform numerical integration over the directions $u$; on the other hand, if $\rho$ is not easily manipulable, we need numerical integration also to obtain the value $w_B(L)$.

5 Results of numerical experiments

In this section we discuss our implementation of the algorithm for the computation of the electrostatic energy of a generic polyhedral object. Then we present numerical experiments, for the case of a homogeneously charged cube and a homogeneously charged parallelepiped. Overall, we tried six different integration methods for the evaluation of the integral (13), and two different implementations of the computation of the kernel (14).

5.1 Setup of numerical experiments

We first implemented the algorithm that computes $K(u)$ in C++ language. We used the C++ class library LEDA (Library of Efficient Data types and Algorithms) [39], available on the internet at the address http://www.mpi-sb.mpg.de. This library contains a routine
that computes the planar decomposition induced by a set of segments, using the sweep-type algorithm described in [38].

The numerical experiment consisted in computing expression (10) using different quadrature schemes for integrating over the directions. We tried the following quadrature schemes:

1. Monte Carlo integration; for this scheme, there exist also theoretical results that relate the behaviour of the error to the geometry of the body [40];

2. Quasi-Monte Carlo integration [41]; FORTRAN routines for generating Halton, Sobol’ and Faure quasi-random sequences of points were taken from the “Collected Algorithms from ACM” [42]; this package is also available on the internet from a variety of sites (e.g. at the URL http://www.math.hkbu.edu.hk/qmc). The experiments were done according to the general framework described in [43];

3. Adaptive multidimensional Gaussian integration; we used routine D01FCF in the NAG library, which is a collection of Fortran routines maintained by the Numerical Algorithms Group [44].

As we will see in the next subsection, the generic implementation, which relies exclusively on standard pieces of software, does not fully exploit the advantages of the new method. In order to obtain better results, we first of all implemented the calculation of the kernel $K(u)$ in C language, using algorithms written on purpose rather than general libraries. Although not highly optimized, this code runs about 40% faster than the previous one.

Moreover, we used a specialized integration algorithm, which is extensively described in [32] and [31]. This algorithm adaptively decomposes the integration domain into subdomains over which the function $K(u)$ is well-behaved; then it performs a standard Gaussian quadrature over each subdomain, summing all the results in the end.

The idea behind the algorithm is that the function $K(u)$ has continuous derivatives in each region where the projection of the polyhedron over $S_u$ is combinatorically the same. While $u$ varies among all the directions, a combinatorial change happens whenever a vertex is projected on the projection of an edge; this corresponds to a certain great circle in the hemisphere of directions $\frac{1}{2}\Omega$, where $K(u)$ may have discontinuous derivatives.
The algorithm tries all the pairs of vertices and edges \((v, e)\) and identifies the great circle which corresponds to the directions \(u\) such that \(v\) is projected onto \(e\) in \(S_u\). The union of these great circles decomposes the hemisphere \(1/2\Omega\) into regions in which the projection of the polyhedron is combinatorically the same. In each such region \(K(u)\) is well-behaved and the Gaussian quadrature succeeds in obtaining a high convergence rate.

The algorithm with the new integration method (which decomposes the domain and then applies Gaussian integration) and the C code for the kernel \(K(u)\) will be denoted with DGQ (Decomposing Gaussian Quadrature). Note that, denoting with \(N\) the number of evaluations of \(K(u)\), the relation \(N \leftrightarrow \text{time}\) depends on the code used for \(K(u)\), while the relation \(N \leftrightarrow \text{accuracy}\) depends on the integration method.

### 5.2 Numerical results

We performed numerical experiments on the computation of the energy of a cube with unit side length, uniformly charged with density \(\rho = 1\). Hummer \[26\] gives an analytic formula for computing the potential of the cube at any point. Integrating this formula over \([0, 1]^3\) with NAG we were able to obtain a very precise reference value for the energy of a unit cube:

\[
E \approx 0.94115632219486 \text{ erg},
\]

with an estimated error of the order of \(10^{-14}\). See Table III to examine the convergence of this integration.

For each of the standard rules of integration discussed in the previous subsection, we carried out 36 different computations, varying the initial rotation of the cube and the seed for the pseudo-random numbers used in the Monte Carlo algorithm. We then computed, for each number \(N\) of function evaluations, the root mean square error, i.e.

\[
\bar{\epsilon}_N = (\sum_j \epsilon_{j,N}^2 / N)^{1/2},
\]

where \(\epsilon_{j,N}\) is the absolute error of the \(j\)-th run after \(N\) function evaluations. Note that, being the value of the integral close to 1, the values \(\epsilon_{j,N}\) represent quite well also the relative error.

The results for the Monte Carlo, Quasi Monte Carlo and NAG integration are shown in Table I. The results obtained with the DGQ method are in Table II. There we show,
Table I: Energy of uniformly charged cube, standard methods

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<th>Faure</th>
<th>Sobol’</th>
<th>NAG</th>
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</table>

Table I: Energy of uniformly charged cube, standard methods

for exponentially increasing values of $N$, the value $-\log_{10} \varepsilon_N$. This number represents the accuracy of the result, in decimal digits, after $N$ evaluations of $K(u)$. The same results are shown graphically in Figure 1, in bilogarithmic scale.

All the computations were done on a Pentium II 200MHz computer. In the tables we also show the time in seconds needed for one run to achieve $N$ evaluations of $K(u)$. The value $\alpha$ at the bottom of each column is the convergence rate of the method. It was obtained by a least-square fitting of the values $\varepsilon_N$, looking for a behaviour like $cN^{-\alpha}$.

In [40] it is shown that the variance of $K(u)$, in the uniform distribution case, is bounded,

Table II: Energy of uniformly charged cube, DGQ method

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<tr>
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so we can expect a convergence rate of $1/2$ from the Monte Carlo method. From the results, it is clear that Monte Carlo method performs exactly as expected. Quasi Monte Carlo methods are superior and have a convergence rate very close to the one predicted by theory (error decreases as $\log^2 N/N$ for two-dimensional integration). The different sequences give a very similar behaviour both in error and convergence rate.

The results of Monte Carlo methods can be compared to those in Morokoff and Caflisch’s work [43]. For a function continuous but not differentiable, defined on a sphere in dimension 2, they exhibit convergence rates between 0.5 (M.C.) and 1.00 (Q.M.C.), with an accuracy of 1.6 digit (M.C.) and 2.85–3.22 digits (Q.M.C.) when $N = 32768$.

The behaviour of the NAG routine is better than that of Monte Carlo method, but the presence of discontinuities in $K(u)$ is a great obstacle in achieving a high convergence rate. Instead, the DGQ method, which prevents discontinuities by subdividing the domain, achieves an error of $10^{-11}$ in only 16 seconds. We stress that this method is general and does not use any property of the cube, so we expect a similar behaviour for any object not stretched in one direction.

Exploiting symmetries

In the integration above we did not take into account the symmetry of the cube when integrating over the directions $u$. However, in real applications, exploiting the symmetry of the objects involved can save a lot of computations. If we do exploit the symmetry of the cube, we can restrict integration to only $1/24$ of the set $\frac{1}{2}\Omega$ (for example in the set $\phi \in [0, \pi/4], z \in [(2 + \tan^2 \phi)^{-1/2}, 1]$). In this way we limit the integration in a region where $K(u)$ is well-behaved, and we do not waste computations for points which give the same value. Although in a similar way, we exploit symmetry better than how the DGQ method does automatically. Note however that in this setting the DGQ method is not applicable.

We carried out numerical experiments in this setting, using the standard integration techniques and the C++ code for $K(u)$. Monte Carlo method was run with 36 different seeds for the generator; for QMC methods we used, in each run, the successive $10^5$ point in the sequence; the NAG routine was run only once, being a completely deterministic algorithm.
The results are shown in Table III. Times of computation are the same as in Table I, although if we use the C code for $K(u)$ we could expect a 40% saving. As a reference, the last column shows the accuracy obtained in the same time of computation when we computed the reference value (15) by 3d adaptive integration of the analytic formula by Hummer.

The analysis shows that the adaptive integrator exploits the new setting achieving a very high accuracy. An error below $10^{-13}$ is the minimum attained, approximately in 16 seconds if we use the C code, even faster than how we obtained the reference value. All three QMC algorithms perform slightly better, in terms of digits of accuracy, but the convergence rates are approximately the same. Monte Carlo method does not seem to gain from the use of symmetry.

**Another example: a parallelepiped**

One can notice that for the cube the integrand function $K(u)$ is very well behaved, it the sense that it does not vary much. In fact, it holds $0.8 \leq 6K(u) \leq 1$ for every direction $u$. 

<table>
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<tr>
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\[ \alpha = 0.56 \quad \alpha = 1.04 \quad \alpha = 1.05 \quad \alpha = 0.98 \quad \alpha = 4.03 \]

Table III: Energy of cube, exploiting symmetries
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<td>2.76</td>
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<td>3.18</td>
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<td>3.20</td>
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<tr>
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<td>1.93</td>
<td>3.60</td>
<td>3.51</td>
<td>3.63</td>
<td>6.18</td>
</tr>
</tbody>
</table>

| $\alpha = 0.51$ | $\alpha = 0.94$ | $\alpha = 0.93$ | $\alpha = 0.95$ | $\alpha = 2.04$ |

Table IV: Energy of parallelepiped, standard methods

This is not the case for general polyhedra, especially if they are stretched in one direction.

In order to better examine this case, we repeated the same numerical experiments taking a $10 \times 1 \times 1$ parallelepiped (with uniform charge density $\rho = 1$). By adapting the formula in [26] we obtained a reference value of 28.52126794 erg for its energy. Notice that in this case the quotient between the maximum and the minimum value of $K(u)$ is 100. Table IV, Table V and Figure 2 show the results, obtained as before by averaging over 36 different runs of the algorithm. The errors shown are the relative errors.

One can see that, as expected, the accuracy is worse than in the case of the cube, even if the asymptotic convergence rates are very similar. Moreover, Monte Carlo and Quasi Monte Carlo methods seem more sensitive to the variation of $K(u)$ than the adaptive integrator. In any case, the DGQ method is much faster than all the others.

Notice that in real applications objects stretched as our parallelepiped are rarely present, so one should expect a behaviour somewhere in-between the cube and the parallelepiped case.
<table>
<thead>
<tr>
<th>( N )</th>
<th>time</th>
<th>DGQ</th>
</tr>
</thead>
<tbody>
<tr>
<td>256</td>
<td>1.0</td>
<td>2.00</td>
</tr>
<tr>
<td>576</td>
<td>2.3</td>
<td>3.08</td>
</tr>
<tr>
<td>1024</td>
<td>4.0</td>
<td>3.63</td>
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<tr>
<td>1600</td>
<td>6.3</td>
<td>3.72</td>
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<tr>
<td>2304</td>
<td>9.1</td>
<td>4.03</td>
</tr>
<tr>
<td>3136</td>
<td>12</td>
<td>4.41</td>
</tr>
<tr>
<td>4096</td>
<td>16</td>
<td>4.82</td>
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<td>5184</td>
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<td>57</td>
<td>7.70</td>
</tr>
<tr>
<td>16384</td>
<td>65</td>
<td>7.80</td>
</tr>
</tbody>
</table>

Table V: Energy of parallelepiped, DGQ method

6 Conclusions

In this paper we have shown that 6-dimensional integrals defining the Coulomb self-energy of a charge distribution and the mutual energy of two distributions can be reduced to 2-dimensional integrals by using integral geometric transformations. This technique is particularly effective for convex polyhedral domains and polynomial distribution of charge, since in this case some auxiliary computation can be done exactly. Preliminary tests on the self-energy of the charged cube, for which reliable reference values are available through an alternative method, show a ratio of precision vs. computational effort comparable to those of other methods in literature aimed at 3-dimensional integrals.
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Figure 1: Energy of uniformly charged cube
Figure 2: Energy of parallelepiped