Nearly uniform sampling of crystal orientations

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June 21, 2018

Abstract

A method is presented for generating nearly uniform distributions of 3D orientations in the presence of symmetry. The method is based on the Thomson problem, which consists in finding the configuration of minimal energy of \(N\) electrons located on a unit sphere—a configuration of high spatial uniformity. Orientations are represented as unit quaternions, which lie on a unit hypersphere in 4D space. Expressions of the electrostatic potential energy and Coulomb’s forces are derived by working in the tangent space of orientation space. Using the forces, orientations are evolved in a conventional gradient-descent optimization until equilibrium. The method is highly versatile as it can generate uniform distributions for any number of orientations and any symmetry, and even allows to prescribe some orientations. For large numbers of orientations, the forces can be computed using only the close neighbourhoods of orientations. Even uniform distributions of as many as \(10^6\) orientations, such as those required for dictionary-based indexing of diffraction patterns, can be generated in reasonable computation times. The presented algorithms are implemented and distributed in the free (open-source) software package Neper.

1 Introduction

Distributions of 3D orientations are routinely used in studies of the physical and mechanical properties of crystalline materials. Examples of experimental studies involve forward simulations of electron backscatter diffraction (EBSD) [1], electron transmission diffraction [2], and high-energy X-ray diffraction [3]. In these studies, simulated diffraction patterns are compared to experimental ones to identify the experimental orientations (or phases), in an approach referred to as “dictionary-based indexing” [2]. Proper indexing requires orientation distributions (for simulating the diffraction patterns) that pave orientation space with a step size of no more than \(1–2^\circ\), which can only be attained by using as many as \(10^6–10^7\) orientations (in the absence of crystal symmetry). For computational efficiency, it is important to minimize this number by distributing orientations as uniformly as possible. Examples of numerical studies in which orientation distributions are used concern simulations of the deformation of polycrystals. Random orientations are usually assigned to the grains, but uniformly-distributed orientations could be used to improve orientation space coverage when analysing the orientation dependency of a mechanical or physical property [4–7], or to reduce the variability of the polycrystal response in homogenization simulations [8]. Such numerical studies typically use polycrystals with \(10^2–10^4\) grains, so that orientation sets are significantly smaller than for dictionary-based indexing.

The general problem of generating uniform distributions of (3D) orientations, or “uniform sampling of orientation space”, has been studied for a long time [9–13] but only rarely in the context
of crystalline materials, which involves symmetries. In the past few years, two methods have been proposed to generate orientation distributions for dictionary-based indexing [1, 14]. Both methods parameterize orientations as unit quaternions to take advantage of the proportionality between the intrinsic distance between unit quaternions and the angle between their corresponding orientations [15]. The problem is then to generate a uniform distribution of points on the unit-quaternion hypersphere. Rosca et al. [1] combine two homochoric projections to transform a regular grid of points in the cube into a regular grid of points on the unit-quaternion hypersphere. Larsen and Schmidt [14] generate a Voronoi tessellation (and its dual, the Delaunay triangulation) by considering the unit quaternions as seed points and combine several optimizations to maximize the minimum distance between any orientation pair. Rosca et al.’s method [1] is direct and therefore very computationally-efficient, and it provides orientation grids that are iso-latitudinal (as needed for the spherical harmonics expansions used in texture-related computations). However, it applies only to specific numbers of orientations ($n^3$), generates orientations that may be relatively close to each other (as equal volumes about orientations do not guarantee equal distances between them), and accounts for crystal symmetry only a posteriori, by rejecting orientations located outside the fundamental region, which leads to undesirable boundary effects. Larsen and Schmidt’s method [14] provides orientation distributions which are effectively more uniform (although not iso-latitudinal), but the method may not behave well for relatively small numbers of orientations due to an “intended use of Euclidean geometries” (i.e., the use of the extrinsic, Euclidean distance in 4D space instead of the intrinsic distance on the unit-quaternion hypersphere) [14]. Finally, it should be noted that both methods [1, 14] involve relatively complex mathematical developments.

In this article, we describe a relatively simple and versatile method that provides improved uniform sampling of crystal orientations. In Section 2, we formulate the method and describe how to solve it. Simplifications are made for the case of large numbers of orientations. In Section 3, we apply the method to several practical cases and compare the results to those of previous methods. In Section 4, we close the article with conclusions. All algorithms described in this article are implemented and distributed in the free (open-source) software package Neper [16].

2 Method

The principle of the method is to adapt the Thomson problem [17] to orientations represented as unit quaternions. The original Thomson problem consists of finding the configuration of minimal energy for a set of electrons located on a unit sphere [17]. It is well-known that this configuration is characterized by a uniform spatial distribution of electrons, for which all electrons occupy about the same surface area of the sphere and exhibit similar distances to their first neighbours. Although formulated back in 1904, the Thomson problem remains open, and several methods have been proposed to solve it, such as gradient-descent optimization [18], global optimization [19], simulated annealing algorithms [20], Monte Carlo algorithms [21, 22], or genetic algorithms [23, 24]. A major difficulty is that the number of local minima grows exponentially with the number of electrons [18, 25]. However, these minima are very close to the global minimum [18,25], so that gradient-descent optimization should provide nearly-optimal configurations. Another difficulty is that the problem becomes computationally-intensive for large numbers of electrons (say, more than 10,000), as all electrons interact with each other. This is not dealt with in most studies (such as [18–24]), which rather focus on finding, and analysing, nearly-optimal configurations for relatively small numbers of electrons (less than 1000), and it will therefore need a specific treatment.

When the Thomson problem is transposed to orientations, several differences arise in its formulation. First, while electrons can interact along chords of the unit sphere (in 3D space), orientations
(represented as unit quaternions) are defined only on the unit-quaternion hypersphere (the 3-sphere in 4D space) and, as such, should interact along its geodesics. Second, in contrast to an electron being a standalone object, an orientation is represented by two (opposite) quaternions or even more (in the presence of crystal symmetry). In this section, we start by reminding the definition and some useful properties of quaternions (Section 2.1). We then present a general formulation for interactions between orientations in the general case and in the presence of crystal symmetry (Section 2.2), and treat the case of large numbers of orientations (Section 2.3). We finally provide a simple and efficient method to solve the problem (Section 2.4).

2.1 Orientations as unit quaternions

Quaternions are 4D vectors that can be used to describe 3D rotations [26]. In fact, there is an homomorphism between the multiplicative group of unit quaternions, SU(2), and the group of rotations in $\mathbb{R}^3$, SO(3) [15]. The space of quaternions, $\mathbb{H}$, is described by the real axis and three imaginary axes along the vectors $i$, $j$, and $k$, so that a quaternion, $q$, can be written as $q = q_0 + q_1 i + q_2 j + q_3 k$ ($q_0$ being the real component and $q_1, q_2, q_3$ being the imaginary components). The base vectors $i$, $j$, and $k$ follow the Hamilton rules for multiplication: $i j = k$, $j k = i$, $k i = j$, $j i = -k$, $k j = -i$, and $i k = -j$. A unit quaternion, $q$, can be defined from a rotation of angle $\theta$ around an axis $r$ ($\|r\| = 1$, where ‘$\|$’ denotes the Euclidean norm) as

$$q = \cos \frac{\theta}{2} + r \sin \frac{\theta}{2}.$$  \hspace{1cm} (1)

Unit quaternions lie on the unit 3-sphere centered at the origin of $\mathbb{H}$, denoted $S^3$, and simply referred to as “unit-quaternion hypersphere” throughout the article. For future reference, we note that the (3D) surface area of $S^3$ is $2 \pi^2$. Moreover, it can be noted that opposite quaternions, $q$ and $-q$, which are antipodal points on $S^3$, correspond to rotations of $(\theta, r)$ and $(\theta + 2 \pi, r)$, respectively, and so represent the same orientation. It is therefore sufficient to consider only positive quaternions, i.e. quaternions of non-negative real parts, to represent orientations (although it is not required in this work). These quaternions occupy the positive hemisphere of $S^3$, $S^3_+$. Two rotations, $q_1$ and $q_2$, both expressed in the reference coordinate system, can be combined into a single rotation by quaternion multiplication as $q_2 q_1$. Quaternion multiplication is associative, distributive, but not commutative. The inverse rotation is given by the conjugate of the original quaternion, which corresponds to inverting the rotation axis (see Equation (1)). Finally, the misorientation between two orientations, $q_1$ and $q_2$, expressed in the reference coordinate system, is simply given by $q_2 q_1^{-1}$.

2.2 Orientation interactions

We formulate the problem of the interaction of orientations represented as electrically-charged quaternions. The aim is to determine the energy of the orientation set and the forces at orientations that derive from it. For simplicity, we assume values of Coulomb’s constant ($k$) and the elementary charge ($e$) equal to 1. We consider a set of two orientations before incorporating crystal symmetry. We then generalize the formulation to a set of $N$ orientations.

2.2.1 Case of a set of two orientations

Let us consider a set of two orientations, $q_i$ and $q_j$, and determine its energy, $E$, and the resulting forces at orientations, $f_i$ and $f_j$ ($f_j = -f_i$), of norm $f$. As pointed out before, $q_i$ and $q_j$ are defined on $S^3$. They therefore interact on $S^3$ (not in its embedding space, $\mathbb{H}$), and so the distances and directions between them are to be measured along geodesics (not chords) of $S^3$; i.e.,
the intrinsic distance is to be used. This is consistent with the fact that the difference between two orientations, measured by the misorientation angle, is proportional to the intrinsic distance between their quaternions. Working on $S^3$ is important to measure distances between orientations properly, especially when the number of orientations is small. It also follows that the forces at orientations are tangential to $S^3$, as can be easily pictured in the 3D case. In other words, the force at an orientation belongs to the tangent space at that orientation. It should therefore be possible to compute it in that space.

The tangent space at an orientation is obtained by logarithm mapping [15], so that the position of $q_j$ in the tangent space at $q_i$, $d_{ij}$, is given by

$$d_{ij} = \ln(q_j q_i^{-1}) = \frac{\theta_{ij}}{2} r_{ij},$$  \hspace{1cm} (2)

where $q_j q_i^{-1}$ corresponds to the misorientation between $q_i$ and $q_j$, and $\theta_{ij}$ and $r_{ij}$ are the corresponding rotation angle and axis (see Section 2.1). It should be noted that even if $q_i$ and $q_j$ are taken as positive quaternions, $q_j q_i^{-1}$ is of arbitrary sign, of real part defined in $[-1, 1]$, and so $\theta_{ij} \in [0, 2\pi]$. $d_{ij}$ is a pure imaginary quaternion, and the space to which it belongs is the 3D ball of radius $\pi$. The geodesics passing through $q_i$ on $S^3$ correspond to straight lines passing through the origin in tangent space, and distances are conserved along these lines [15]. The distance between $q_i$ and $q_j$ (either on $S^3$ or in tangent space), $d_{ij}$, is

$$d_{ij} = \|d_{ij}\| = \frac{\theta_{ij}}{2},$$  \hspace{1cm} (3)

and the (signed) direction between $q_i$ and $q_j$ is equal to $r_{ij}$. Since $\theta_{ij} \in [0, 2\pi]$, we have $d_{ij} \in [0, \pi]$. As the orientation represented by $q_j$ can also be represented by $-q_j$, $q_i$ interacts with both $q_j$ and $-q_j$. We may compute the position of $-q_j$ in the tangent space at $q_i$, $d'_{ij}$, as

$$d'_{ij} = \ln(-q_j q_i^{-1}) = \frac{\theta'_{ij}}{2} r'_{ij}.$$  \hspace{1cm} (4)

Noting that $d_{ij}$ and $d'_{ij}$ are mapping opposite quaternions, and using Equation (1), it is easy to show that $\theta'_{ij} = 2\pi - \theta_{ij}$ and $r'_{ij} = -r_{ij}$. Therefore, $d'_{ij}$ can be written as

$$d'_{ij} = -\frac{2\pi - \theta_{ij}}{2} r_{ij},$$  \hspace{1cm} (5)

and it follows that (also using Equation (3))

$$d'_{ij} = \|d'_{ij}\| = \frac{2\pi - \theta_{ij}}{2} = \pi - d_{ij}.$$  \hspace{1cm} (6)

As $d_{ij} \in [0, \pi]$, we also have $d'_{ij} \in [0, \pi]$. It can finally be noted that $d_{ij}$ and $d'_{ij}$ are parallel but point in opposite directions, and that if one of $d_{ij}$ or $d'_{ij}$ belongs to the 3D ball of radius $\pi/2$, the other belongs to the 3D spherical shell of inner radius $\pi/2$ and outer radius $\pi$, as illustrated on Figure 1(a). The same calculations could be carried out by considering $-q_i$ instead of $q_i$, which would provide the same distances ($d_{ij}$ and $\pi - d_{ij}$). The total energy of the set of two orientations ($E$) results from the interactions between the four pairs $(q_i, q_j)$, $(q_i, -q_j)$, $(-q_i, q_j)$ and $(-q_i, -q_j)$, and is given by

$$E = 2 \left( \frac{1}{d_{ij}} + \frac{1}{\pi - d_{ij}} \right).$$  \hspace{1cm} (7)
The force at \( q_i \), \( f_i \), derives from the energy that \( q_i \) forms with \( q_j \) and \( -q_j \), which is equal to \( E/2 \). Moreover, as the force is repulsive, \( f_i \) is opposite to \( r_{ij} \). So, the force at \( q_i \) can be written in scalar and vector forms as

\[
f = -\frac{d(E/2)}{dd_{ij}}, \quad f_i = -f r_{ij}. \tag{8}
\]

Substituting \( E \) by its expression (Equation (7)) and applying the derivative, we finally obtain

\[
f = \frac{1}{d_{ij}^2} - \frac{1}{(\pi-d_{ij})^2}, \quad f_i = -\left(\frac{1}{d_{ij}^2} - \frac{1}{(\pi-d_{ij})^2}\right) r_{ij}. \tag{9}
\]

Figure 1(b) shows the evolutions of \( E \) and \( f \) as a function of \( d_{ij} \). As expected, a minimal value of \( E \) and a zero value of \( f \) are obtained for \( d_{ij} = \pi/2 \) (\( \theta_{ij} = \pi \)), i.e. when \( q_i \) and \( q_j \) represent opposite orientations (there actually is a family of quaternions \( q_j \) representing orientations opposite to \( q_i \), which corresponds to the sphere of radius \( d_{ij} = \pi/2 \) on Figure 1(a) (\( \theta_{ij} = \pi \), arbitrary \( r_{ij} \)).

Finally, it should be noted that Equations (7) and (9) differ from the corresponding concepts known from electrostatics in several ways. First, antipodal symmetry leads to an equilibrium configuration at a finite distance, where forces vanish. Second, and strictly speaking, the force equilibrium between the two orientations, \( f_j = -f_i \), can only be written under the concept of parallel transport. Indeed, \( f_i \) and \( f_j \) have the same magnitude (\( f \)), but they belong to different tangent spaces and so have different directions in \( H \). Only when \( f_j \) is transported from \( q_j \) to \( q_i \) (along the geodesics between \( q_j \) and \( q_i \)) does the standard expression of force equilibrium hold true. This will, however, not be used in the following since the forces at orientations will always be computed in their respective tangent spaces.
2.2.2 Crystal symmetry

In the presence of crystal symmetry, an orientation, represented by quaternion \( q \), can be described by several, equivalent quaternions, \( q^k \), \( k = 1, \ldots, n_c \), given by

\[
q^k = q u^k,
\]

where \( u^k \) is a symmetry operator. For cubic crystal symmetry, \( n_c = 24 \) and \( u^k \) can represent the identity, the three rotations of \( \pi/2 \) about each of the three \( \langle 100 \rangle \) crystal directions, the rotation of \( \pi \) about each of the six \( \langle 110 \rangle \) crystal directions, and the two rotations of \( 2\pi/3 \) about each of the four \( \langle 111 \rangle \) crystal directions. To compute \( E \) and \( f_i \), all equivalent orientations must be taken into account. Let us consider the interaction between \( q_i \) and \( q_j^k \). The misorientation axis and angle between \( q_i \) and \( q_j^k \) are denoted as \( r_{ij}^k \) and \( \theta_{ij}^k \), respectively. \( q_j^k \) can be substituted by \( q_j^{k'} \) in Equations (2) and (3) to compute the distance between \( q_i \) and \( q_j^{k'} \), \( d_{ij}^{k'} \). The total energy of the pairs formed by \( q_i \) and all \( q_j^{k'} \)s (and their negatives) can simply be written as the sum of the energies of all orientation pairs (which are of the form of Equation (7)). It should then be noted that, due to symmetry considerations, the same result would be obtained by considering \( q_i^k (k = 1, \ldots, n_c) \) instead of \( q_i \). The total energy therefore is

\[
E = 2 n_c \sum_{k=1}^{n_c} \left( \frac{1}{d_{ij}^k} + \frac{1}{\pi - d_{ij}^k} \right).
\]

The net force at an orientation, \( f_i \), can be written as the sum of the forces between the orientation \( (q_i) \) and all other orientations \( (q_j^k) \) (which are of the form of Equation (9)):

\[
f_i = - \sum_{k=1}^{n_c} \left( \frac{1}{d_{ij}^k} - \frac{1}{\pi - d_{ij}^k} \right) r_{ij}^k.
\]

2.2.3 General case

Let us now consider a set of \( N \) orientations, \( q_i, i = 1, \ldots, N \), with crystal symmetry. The total energy of the orientation set \( (E) \) can be written as the sum of the energies of all orientation pairs (see Equation (11)):

\[
E = 2 n_c \sum_{i=1}^{N} \sum_{j=i+1}^{N} \sum_{k=1}^{n_c} \left( \frac{1}{d_{ij}^k} + \frac{1}{\pi - d_{ij}^k} \right).
\]

The net force at an orientation \( (f_i) \) can be written as the sum of the forces between the orientation and all other orientations (see Equation (12)):

\[
f_i = - \sum_{j=1, j \neq i}^{N} \sum_{k=1}^{n_c} \left( \frac{1}{d_{ij}^k} - \frac{1}{\pi - d_{ij}^k} \right) r_{ij}^k.
\]

As expected from the original Thomson problem [17] and as will be seen in Section 3, a minimal value of the energy \( (E) \) and zero values of the forces \( (f_i) \) are obtained when orientations are uniformly distributed on \( S^3 \).
2.3 Case of a large number of orientations

As the energy and the forces (provided by Equations (13) and (14)) depend on all orientations pairs, computing them can be relatively long \((O(N^2))\) when the number of orientations \((N)\) is large. However, it should first be noted that only the forces (not the energy) must be computed during the resolution (see Section 2.4), and second that the forces should be mostly driven by short-distance interactions, as the (many) long-distance interactions yield low forces that tend to cancel each other out. It should therefore be sufficient to compute the forces in close neighbourhoods of the orientations. In contrast to what was done in Section 2.2, it is here useful to write the expression of the forces by representing all quaternions (positive and negative) equivalently, so that Equation (14) becomes

\[
 f_i = -\sum_{q^\prime \in \{q^k_j, -q^k_j\}, j \neq i} \frac{1}{d(q_i, q'^\prime)^2} r(q_i, q'),
\]

where \(d(q_i, q)\) and \(r(q_i, q)\) denote the distance and the misorientation axis between \(q_i\) and \(q\), respectively (and correspond to \(d_{ij}\) and \(r_{ij}\) for \(q \in \{q^k_j\}\), and to \(d'_{ij}\) and \(r'_{ij}\) for \(q \in \{-q^k_j\}\), respectively). An approximate value of \(f_i\) can then be obtained by considering only the quaternions located in the neighbourhood of \(q_i\):

\[
 f_i \simeq -\sum_{q \in \{q^k_j, -q^k_j\}, d(q_i, q) \leq \delta d_r} \frac{1}{d(q_i, q)^2} r(q_i, q),
\]

where the neighbourhood corresponds to an hyperspherical cap (on \(S^3\)) of radius \(\delta d_r\) and centre \(q_i\). \(\delta\) is a multiplicative factor (to be determined in Section 3.3), and \(d_r\) is the so-called “average orientation radius” and corresponds to the radius of an hyperspherical cap of surface area, \(S\), defined as

\[
 S = \frac{\pi^2}{N^*},
\]

where \(N^* = N n_c\). \(S\) is related to the angular radius of the corresponding hyperspherical cap, \(\theta_r\), by [15]

\[
 S = \pi (\theta_r - \sin \theta_r),
\]

Knowing \(N\) and \(n_c\), \(\theta_r\) can be computed using Equations (17) and (18), and \(d_r\) is given by \(\theta_r/2\). Knowing \(\delta d_r\), neighbours are rapidly found using a kD-tree technique (in \(O(N \log N)\)) [27].

2.4 Resolution

The energy of an orientation distribution and the resulting forces being known (Equations (13) and (14)), the energy can be minimized using a conventional gradient-descent method, which iteratively evolves the orientations along (and proportionally to) their respective forces.

2.4.1 General formulation

During an iteration \((l)\), each orientation evolves from its initial position, \(q_i^{(l-1)}\), to its final position, \(q_i^{(l)}\). The rotation during the iteration, \(\Delta q_i^{(l-1)}\), is computed from the force at the beginning of the iteration, \(f_i^{(l-1)}\), as

\[
 \Delta q_i^{(l-1)} = \alpha^{(l-1)} f_i^{(l-1)},
\]
where \( \alpha^{(l-1)} \) is the step size of the iteration \((l)\) and controls the degree of evolution of the orientations. \( \alpha^{(l-1)} \) is the same for all orientations and is computed as described in Section 2.4.2. The orientation evolution during the iteration (i.e., the calculation of \( q_i^{(l)} \) from \( q_i^{(l-1)} \) and \( \Delta q_i^{(l-1)} \)) can be expressed as a simple additive composition, although it needs more discussion. It should be recalled that while \( q_i \) is a unit quaternion, i.e. a point on \( \mathbb{S}^3 \), \( f_i \) and therefore \( \Delta q_i \) belongs to the tangent space at \( q_i \). This space is the space of pure imaginary quaternions and is centred at the origin of the space of unit quaternions, \((1, 0, 0, 0)\). \( \Delta q_i \) is therefore tangential to \( \mathbb{S}^3 \) at \((1, 0, 0, 0)\), not at \( q_i \). An additive composition requires \( \Delta q_i \) to be rotated to \( q_i \) (through parallel transport), which leads to \( \Delta q_i^* \):

\[
\Delta q_i^* = \Delta q_i q_i.
\]

The orientation evolution during an iteration can then be written as

\[
q_i^{(l)} = q_i^{(l-1)} + \Delta q_i^{(l-1)};
\]

i.e., the additive composition is subjected to a normalization (represented by ‘\( \sim \)’). Normalization is necessary since, during the iteration, and as \( \Delta q_i \) is tangential to \( \mathbb{S}^3 \), \( q_i \) moves along \( \mathbb{S}^3 \) but also deviates from it to second order.

### 2.4.2 Step size and termination criterion

The step size \((\alpha^{(l-1)})\) used in Equation (19) must be set appropriately throughout optimization, so that the fewest possible iterations are needed. This could be done by determining, at each iteration and through a line search, the value of \( \alpha \) that most decreases the energy. However, this would require many (expensive) computations of \( E \) (one for each value of \( \alpha \) tested). Instead, we adopt some assumptions to compute values of \( \alpha^{(l-1)} \) from information known \( \text{a priori} \) or from the previous iteration.

The value of \( \alpha \) at the first iteration, \( \alpha^{(0)} \), is defined so that it provides a maximal decrease of \( E \), i.e. a minimal value of \( E^{(1)} \), for a typical random orientation distribution. A minimal value of \( E^{(1)} \) is obtained for specific amplitudes of the rotations, \( \| \Delta q_i^{(0)} \| \) or, equivalently (as all rotations evolve according to Equation (21)), for a specific value of their average, \( \langle \| \Delta q_i^{(0)} \| \rangle \). The average amplitude of the rotations \( \langle \| \Delta q_i^{(0)} \| \rangle \) can be expressed as a fraction, \( \beta \), of the average orientation radius \((d_r)\). This can be formulated as

\[
\langle \| \Delta q_i^{(0)} \| \rangle = \beta d_r \quad \text{with} \quad \langle \| \Delta q_i^{(0)} \| \rangle \text{ such that } E^{(1)} \text{ minimum.}
\]

By combining Equations (19) and (22), \( \alpha^{(0)} \) can be written as

\[
\alpha^{(0)} = \frac{\beta d_r}{\langle \| f_i^{(0)} \| \rangle},
\]

where \( \langle \| f_i^{(0)} \| \rangle \) corresponds to the average of the forces at orientations in the initial configuration. It should be noted that \( f_i^{(0)} \) is of the form \( 1/d_{ij}^2 \) (neglecting the term in \( 1/(\pi - d_{ij})^2 \) in Equation (9), as in Section 2.3) and therefore depends mostly on interactions within the close neighbourhood of \( q_i^{(0)} \). The distances to the orientations within the neighbourhood \((d_{ij})\) are proportional to \( d_r \), and so \( \langle \| f_i^{(0)} \| \rangle \) is inversely proportional to \( 1/d_r^2 \):

\[
\langle \| f_i^{(0)} \| \rangle = \frac{\gamma}{d_r^2}.
\]
By combining Equations (23) and (24), $\alpha^{(0)}$ can finally be written as

$$\alpha^{(0)} = \frac{\beta}{\gamma} d_r^3. \tag{25}$$

It is then interesting to determine the values of $\beta$ and $\gamma$ independently, which can be done using, as values of $q_i^{(0)}$, a random distribution of $10^5$ orientations (considered sufficiently large to get representative results). The value of $\beta$, as defined in Equation (22), can be determined by a line search (see Figure 2), which provides $\beta = 0.8$. In other words, at the first iteration, a maximal decrease of the energy ($E$) is obtained when orientations are rotated in average by 80% of the average orientation radius ($d_r$). The value of $\gamma$, as defined in Equation (24), can be computed directly, which provides $\gamma = 3.4$.

The values of $\alpha$ at the following iterations are determined using the Barzilai-Borwein method [28]. In general terms, the Barzilai-Borwein method [28] assumes a quadratic form of the minimized function and computes $\alpha$ from the evolutions, during the previous iteration, of the solution vector and of the gradient of the minimized function. In this work, the minimized function is the energy ($E$), the solution vector corresponds to the $q_i$ values and its evolution to the $\Delta q_i$ values, and the gradient of the minimized function corresponds to the forces ($f_i$) and its evolution to the $\Delta f_i$ values. The Barzilai-Borwein method provides two possible values of $\alpha^{(l-1)}$ (which result from two different least-square problems) [28]:

$$\alpha_1^{(l-1)} = \frac{\sum_{i=1}^{N} (\Delta q_i^{(l-2)} \cdot \Delta f_i^{(l-2)})}{\sum_{i=1}^{N} (\Delta f_i^{(l-2)} \cdot \Delta f_i^{(l-2)})},$$

and

$$\alpha_2^{(l-1)} = \frac{\sum_{i=1}^{N} (\Delta q_i^{(l-2)} \cdot \Delta q_i^{(l-2)})}{\sum_{i=1}^{N} (\Delta q_i^{(l-2)} \cdot \Delta f_i^{(l-2)})}.$$

(26)

Which of $\alpha_1^{(l-1)}$ and $\alpha_2^{(l-1)}$ is the most efficient in decreasing $E$ is unknown a priori, and a common practice is to alternate between them during optimization. In this work, we use $\alpha_1^{(l-1)}$ on odd iterations and $\alpha_2^{(l-1)}$ on even iterations.

Finally, the optimization terminates based on a relative error on the forces (a residual):

$$\sqrt{\frac{\sum_{i=1}^{N} ||f_i^{(l)}||^2}{\sum_{i=1}^{N} ||f_i^{(0)}||^2}} < \varepsilon_r. \tag{27}$$

In this article (and except mentioned otherwise), we use $\varepsilon_r = 10^{-3}$, which provides a good compromise between accuracy and computation time.

### 3 Results

The method is validated and applied to several practical cases. In Section 3.1, we describe how the uniformity of an orientation distribution is measured. In Section 3.2, we analyse the method in
3.1 Methodology

The uniformity of orientation distributions is analysed using two metrics. The first metric is the orientation distribution function (ODF), $f(q)$, and we will focus on its standard deviation, $\sigma_f$. The ODF ($f(q)$) is computed using a discrete method, for which each orientation ($q_i$) (and its symmetry equivalents ($q_k^i$)) is represented by a continuous, Bingham distribution (which acts as a smoothing kernel) [15]. The ODF ($f(q)$) can therefore be written as

$$f(q) = \frac{1}{N n_c} \sum_{i=1}^{N} \sum_{k=1}^{n_c} \left( _1F_1 \left[ \frac{1}{2}, 2, K \right] \right)^{-1} \exp \left( w^k_i U K U^t w^k_i \right)$$

with $$w^k_i = q^k_i q^{-1},$$

where $U$ is an orthonormal $4 \times 4$ matrix, $K$ is a diagonal matrix describing the shape of the distributions, and $_1F_1$ is the hypergeometric function with matrix argument normalizing the density of the Bingham distribution. In this work, we consider isotropic Bingham distributions of (unidirectional) standard deviation equal to $d_\tau$, as obtained using $U = I_4$, $K_{11} = 2/d_\tau^2$, and $K_{22} = K_{33} = K_{44} = 0$. Considering $d_\tau$ as the standard deviation provides a level of smoothing that is independent of the number of orientations ($N^*$).

Different orientation parameterizations can be used to visualize the ODF. In the absence of crystal symmetry, the homochoric parameterization [1], $x_h$, is used:

$$x_h = r \left( \frac{3}{4} (\theta - \sin \theta) \right) \frac{1}{2},$$

which maps orientation space into the 3D ball of radius $(3 \pi/4)\frac{1}{2}$ (as will be seen in Figure 3). In the presence of crystal symmetry, it is sufficient to represent the ODF in the fundamental region of
orientation space. The Rodrigues parameterization, \( x_R \), is used:

\[
x_R = r \tan \frac{\theta}{2},
\]

which maps the fundamental region of orientation space into a convex polyhedron whose size and shape depend on the crystal symmetry: for cubic crystal symmetry, the polyhedron is a truncated cube, while for hexagonal crystal symmetry, the polyhedron is a dodecagonal prism (as will be seen in Figure 5). The more uniform an orientation distribution, the closer to 1 its ODF \( f(q) \), and the lower \( \sigma_f \). \( \sigma_f \) has a value of 0.3 for a typical random orientation distribution and a minimal value of 0 for a perfectly uniform orientation distribution. It should however be noted that, as the ODF is computed from a finite number of orientations, \( \sigma_f \) will never reach the minimal value of 0, but instead a slightly higher, “residual” value that is independent of \( N^* \) (because of the chosen dependence of \( d_r \) on \( N^* \)). The optimal covering in a local area should be the b.c.c. lattice since this is the best known covering in \( \mathbb{R}^3 \) \cite{14}; the corresponding value of \( \sigma_f \) is 0.0021, which can be considered as an absolute minimal value of \( \sigma_f \).

The second metric is the function corresponding to the misorientation between an arbitrary orientation and the nearest orientation of the orientation distribution, \( \theta_m(q) \), and we will focus on properties such as its average or maximum. Unlike \( f(q) \), which is a fundamental and widely-used metric, \( \theta_m(q) \) is relevant specifically in the context of dictionary-based indexing. Although the two metrics are related, they are not strictly equivalent as \( f(q) \) considers that orientations are represented by a kernel (and therefore represent a small region) while \( \theta_m(q) \) depends only on the closest orientation. It results that both metrics may not evolve in the same way, as will be seen in the following.

3.2 Orientation evolution during optimization

The evolution of a distribution of 1000 orientations during optimization is illustrated on Figure 3 (no symmetry). On Figure 3(a), it can be seen that the energy \( (E) \) shows a rapid decrease at the beginning of optimization and then saturates. The energy decreases only from \( 7.75 \times 10^5 \) to \( 7.69 \times 10^5 \) but, at the same time, the ODF standard deviation \( (\sigma_f) \) decreases significantly, from 0.29 down to 0.007. This reveals that the local arrangement of the orientations or, more specifically, its degree of uniformity (which is related to \( \sigma_f \)), represents only a small fraction of the energy (all the more that \( N \) is large). Still, minimizing the energy appears to be very efficient in decreasing \( \sigma_f \). Figure 3(b) provides the ODF at successive iterations, and it can be seen how the ODF, which initially shows large variations, becomes more and more uniform during optimization to eventually reach values very close to 1. On Figure 3, the ODF is shown only on the surface of the homochoric region, but this should not be considered a limitation since surface values are representative of interior values; slice views will be provided for the comparison with other methods (see Figure 7).

3.3 Influence of the local calculation of the forces

The influence of restraining the force calculation to a close neighbourhood (as described in Section 2.3) on the resulting ODF standard deviation \( (\sigma_f) \) can be analysed. The aim is to determine how small the neighbourhood can be (minimal value of \( \delta \)) while still providing a value of \( \sigma_f \) similar to the nominal value. The procedure is as follows: distributions of \( 10^5 \) orientations (no symmetry) are generated for increasing values of \( \delta \), and their respective ODF standard deviations \( (\sigma_f) \) are computed. The evolution of \( \sigma_f \) as a function of \( \delta \) is plotted on Figure 4 for two values of the residual, \( \varepsilon_r = 10^{-3} \) (the default) and \( \varepsilon_r = 10^{-4} \). For both values, and as expected, \( \sigma_f \) tends to decrease as \( \delta \) increases. For \( \varepsilon_r = 10^{-3} \), \( \sigma_f \) becomes close to the nominal value (and lower than 0.01).
Figure 3: Generation of a uniform distribution of 1000 orientations (no symmetry). (a) Evolution of the energy ($E$) and the ODF standard deviation ($\sigma_f$). The dashed lines represent the asymptotic solutions (1000 iterations). (b) Orientation distribution functions shown on the homochoric projection ball at iterations 0 (initial solution), 1, 2, 5, 34 (final solution) and 1000 (asymptotic solution). Note how the evolution of $\sigma_f$ is non monotonic (the same applies to $E$), which is typical from the Barzilai-Borwein method.
for $\delta \approx 20$. For larger values of $\delta$, fluctuations of $\sigma_f$ appear, which are caused by the variations of the residual as it approaches $\varepsilon_r$ (at the end of optimization). As can be seen on Figure 4, $\varepsilon_r = 10^{-4}$ leads to lower values of $\sigma_f$ (and no apparent fluctuations), albeit at the cost of longer computations and larger values of $\delta$ to approach the nominal value. A value of $\delta$ of 20 corresponds to a number of 7500 orientations in the neighbourhood, and so restraining the force calculation to close neighbourhoods (with $\delta = 20$) reduces computation time for $2N^* > 7500$.

### 3.4 Practical applications

The method is first applied to the generation of uniform distributions of 1000 orientations with cubic and hexagonal crystal symmetries. The ODFs corresponding to the initial, random orientation distributions and to the final, uniform orientation distributions are shown on Figure 5. Similarly to the case of triclinic symmetry (Figure 3(b)), the ODFs corresponding to the uniform orientation distributions exhibit significantly lower standard deviations ($\sigma_f$). During the simulation, $\sigma_f$ actually decreases from 0.27 down to 0.007 for cubic symmetry and from 0.320 down to 0.007 for hexagonal crystal symmetry.

The method can be applied to arbitrary numbers of orientations ($N$). Typical computation times are reported on Table 1 for several values of $N$ and several crystal symmetries. Figure 6 provides the values of $\sigma_f$ for $N$ ranging from $10^2$ to $10^6$ in the case of cubic crystal symmetry, for the present method and for previous methods [1,14]. It appears that the present method leads to a value of $\sigma_f$ of about 0.007 independently of $N$, which is significantly lower than those obtained by previous methods [1,14]. More specifically, Roșca et al.’s method [1] provides relatively low values of $\sigma_f$ only for high values of $N$, but these values are still 10–20 times higher than for other methods, while Larsen and Schmidt’s method [14] provides a value of 0.024 independently of $N$. Figure 6 also shows that, unlike the present method, previous methods apply only to specific values of $N$ (those available online, for Larsen and Schmidt’s method [14]). A closer comparison can be made between the different methods by inspecting their ODFs, as illustrated on Figure 7 for $N = 9218$ ($N = 9077$ for Roșca et al. [1]). It can be seen that, unlike other methods, Roșca et al.’s method [1] leads to
strong variations of the ODF near the surface of the fundamental region. This results from the fact that crystal symmetry is only taken into account \textit{a posteriori}, by discarding orientations located outside the fundamental region [1]. As for Larsen and Schmidt’s method [14], it is confirmed that the ODF shows stronger variations compared to the present method.

The different methods can also be compared in terms of the distribution of the misorientations to the nearest orientation ($\theta_m(q)$), as illustrated on Figure 8. A distribution of 262,144 orientations (259,385 orientations for Roşca et al.’s method [1]) with cubic crystal symmetry is considered, and the frequency distribution of $\theta_m(q)$ was computed from the values at $10^8$ arbitrary orientations. It can be seen that the present method and Larsen and Schmidt’s method [14] generate similar distributions, with an average value of $\sim 0.63^\circ$, which is lower than that of Roşca et al.’s method [1], $0.67^\circ$. Larger differences arise for the maximal values: $1.84^\circ$ for Roşca et al.’s method [1], $0.99^\circ$ for Larsen and Schmidt’s method [14], and $1.17^\circ$ for the present method. As a matter of fact, 0.2\% of the values of $\theta_m$ exceeds $0.99^\circ$ for the present method, while 4.5\% do for Roşca et al.’s method [1]. The fact that Larsen and Schmidt’s method [14] produces a lower maximal value of $\theta_m(q)$ (but a higher value of $\sigma_f$) can be attributed to its being specifically designed for that purpose. Actually, the Thomson problem solved in this work and the packing problem solved in Larsen and Schmidt’s method [14] are two specializations of the generalized Thomson problem of minimizing a Riesz energy of the form $\sum_{i\neq j} 1/d_{ij}^s$ [29]. For the Thomson problem, $s = 1$ and the energy depends on the interaction between all pairs, while for the packing problem, $s = \infty$ and only the one orientation pair with the smallest distance matters [29]. This explains why the present method provides better results on $\sigma_f$ (which depends on interactions within a neighbourhood) while Larsen and Schmidt’s method [14] provides better results on the maximum value of $\theta_m(q)$ (which is local).

The improvement provided by a uniform orientation distribution (compared to a random orientation distribution) can finally be analysed from a different perspective: instead of determining how $\sigma_f$ can be decreased (for a fixed $N$), we can determine how $N$ can be decreased (for a fixed $\sigma_f$), as illustrated on Figure 9. This corresponds to looking at how the number of orientations can be decreased while still getting the same “coverage” of orientation space (measured by $\sigma_f$). This can be done by computing the ODFs slightly differently than previously, using the same smoothing kernel for all values of $N$. We can then analyse how $\sigma_f$ changes as a function of $N$, for random and uniform orientation distributions. On Figure 9, it can be seen that, in the case of cubic crystal symmetry, a uniform distribution of 130 orientations has the same value of $\sigma_f$ as a random distribution of 1000 orientations.

A last application concerns the generation of a uniform orientation distribution including prescribed orientations. This is desirable in some applications, for example to analyse the behaviour of specific (experimental) orientations [5, 6]. Figure 10 shows a distribution of 1000 orientations among which 82 orientations were positioned along the $X_i \parallel \langle 100 \rangle$ and $X_i \parallel \langle 110 \rangle$ orientation fibres. It can be seen that, despite the constraints imposed by the prescribed orientations, the final orientation distribution is as uniform as in the standard case (see Figure 7(c)), which indicates that the unconstrained orientations have adapted to the prescribed orientations during optimization.

4 Conclusion

By transposing the well-known Thomson problem for the equilibrium distribution of electrons on a unit sphere to orientations represented as unit quaternions, we have devised a relatively simple, versatile, and efficient method for generating uniform orientation distributions. The key points of the method are the following:

1. Owing to the fact that 3D orientations represented as unit quaternions are defined on $S^3$ (not
Figure 5: Comparison of a random distribution (initial solution) and a uniform distribution (final solution) of 1000 orientations in the presence of crystal symmetry. ODFs are shown in the fundamental regions of Rodrigues space. (a) Cubic crystal symmetry and (b) hexagonal crystal symmetry.

Table 1: Computation times for generating uniform distributions for several numbers of orientations ($N$) and crystal symmetries. For $N \geq 10^5$, forces are computed in neighbourhoods ($\delta = 20$). Computation times are averaged over 50 simulations (starting from different random orientation distributions). Simulations are run on 10 cores (20 threads) of an Intel® Xeon® CPU E5-2660 v3 processor. Note that the computation times are similar for all symmetries for $N = 10^6$, which is a consequence of the force computation in close neighbourhoods.

<table>
<thead>
<tr>
<th>$N$</th>
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<th>hexagonal</th>
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<td>$10^3$</td>
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<tr>
<td>$10^4$</td>
<td>4min</td>
<td>36min</td>
<td>22min</td>
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<tr>
<td>$10^5$</td>
<td>17min</td>
<td>45min</td>
<td>31min</td>
</tr>
<tr>
<td>$10^6$</td>
<td>3h30min</td>
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</tbody>
</table>

15
in its embedding space, \( \mathbb{H} \), they were considered to interact along its geodesics, so that the forces at orientations could be elegantly written in the tangent space at orientations. This both allows a simple calculation and avoids biases, especially for small numbers of orientations.

2. The forces at orientations, being tangential to \( S^3 \), provide evolution paths and rates (for the orientations) toward states of lower energies, which was used in a conventional gradient-descent optimization. Although this corresponds to local optimization and does not ensure a global minimum, the resulting orientation distributions are highly uniform.

3. In the exact formulation, the force at an orientation depends on its interaction with all other orientations, so that large numbers of orientations \( (N) \) leads to very high computation times \( (O(N^2)) \). However, we showed that for more than about 7500 orientations, the forces can be computed in close neighbourhoods of the orientations, as forces mostly depend on short-distance interactions. This significantly reduces computation time \( (O(N \log N)) \), so that large values of \( N \) can be used (e.g. \( 10^6 \)).

4. Crystal symmetry can be included naturally, by representing each orientation by all its symmetrically-equivalent quaternions. Similarly, orientations that are desired in the distribution can be prescribed.

5. An efficient resolution and implementation enables to generate uniform distributions of as many as \( 10^6 \) orientations within reasonable computation time. A parallel (MPI) implementation could be used to reduced computation time further.

The uniformity of the orientation distributions was analysed using two metrics: the orientation distribution function \( (f(q)) \), and the distributions of the misorientations to the nearest orientation \( (\theta_m(q)) \), both of which are of practical importance for crystalline materials. The present method provides highly uniform orientation distributions in terms of the standard deviation of \( f(q) (\sigma_f) \) and the average value of \( \theta_m(q) \). Uniform distributions for any number of orientations (from only a few to \( 10^6 \) or more) and any crystal symmetry can be generated and used routinely.
Figure 7: ODFs provided by different methods, for ∼9200 orientations and cubic crystal symmetry: (a) Roșca et al. [1] (9077 orientations), (b) Larsen and Schmidt [14] (9218 orientations) and (c) this work (9218 orientations). ODFs are shown as surface and slice views. Note that the colour scale has been reduced to [0.8, 1.2] compared to Figures 3 and 5.
Figure 8: Frequency distributions of the misorientations to the nearest orientation ($\theta_m(q)$) in the case of cubic crystal symmetry, as obtained by different methods: Cubochoric after Roșca et al. [1], CRO after Larsen and Schmidt [14], and Thomson (this work). $N = 259,385$ for Cubochoric (after Roșca et al.’s method [1]), and $N = 262,144$ for CRO (after Larsen and Schmidt [14]) and Thomson (this work). Average values are: 0.66°, 0.63° and 0.63°, for the three methods, respectively, and maximal values are: 1.84° (out of scale), 0.99°, and 1.17°, for the three methods, respectively. The cubochoric distribution has been scaled to correct for its slightly smaller number of orientations.

Figure 9: Comparison of the orientation space coverages (measured by $\sigma_f$) obtained for random and uniform orientation distributions, in the case of cubic crystal symmetry. A fixed smoothing kernel is used (which corresponds to the nominal kernel for 1000 orientations). The dashed lines indicate how a uniform orientation distribution of 130 orientations provides the same orientation space coverage as a random orientation distribution of 1000 orientations.
Figure 10: Generation of a uniform orientation distribution in the presence of prescribed orientations. Case of 1000 orientations among which 82 are prescribed, which are located along the $X_i \parallel (100)$ and $X_i \parallel (110)$ fibres. The prescribed orientations are shown by blue dots and the color field represents the ODF. Cubic crystal symmetry is considered. ODFs are shown as surface and slice views. Note that the ODF is as uniform as in the standard case (Figure 5(a)).

5 Acknowledgements

One of the authors (R.Q.) wish to thank Adam Morawiec for early discussions on the subject. The authors wish to thank Adam Morawiec, as well as the two anonymous reviewers, for their thorough comments on the manuscript. Finally, the authors would like to dedicate this article to the memory of Julian Driver.

References


