

THREE- AND FOUR-DIMENSIONAL K -OPTIMAL LATTICE RULES OF MODERATE TRIGONOMETRIC DEGREE

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ABSTRACT. A systematic search for optimal lattice rules of specified trigonometric degree d over the hypercube $[0, 1]^s$ has been undertaken. The search is restricted to a population $K(s, \delta)$ of lattice rules $Q(\Lambda)$. This includes those where the dual lattice Λ^\perp may be generated by s points \mathbf{h} for each of which $|\mathbf{h}| = \delta = d + 1$. The underlying theory, which suggests that such a restriction might be helpful, is presented. The general character of the search is described, and, for $s = 3$, $d \leq 29$ and $s = 4$, $d \leq 23$, a list of K -optimal rules is given. It is not known whether these are also optimal rules in the general sense; this matter is discussed.

1. INTRODUCTION

We consider cubature rules for $[0, 1]^s$ of trigonometric degree d . Such a rule integrates correctly all s -dimensional trigonometric polynomials of degree d . Specifically, it integrates $\exp(2\pi i \mathbf{h} \cdot \mathbf{x})$ correctly for all $\mathbf{h} := (h_1, h_2, \dots, h_s) \in \mathbb{Z}^s$ that satisfy $|\mathbf{h}| := \sum_{k=1}^s |h_k| \leq d$. Lattice rules have played a significant role in the development of this area. For background information of a general nature on lattice rules, we refer to [SJ94] and to [CS96] for lattice and other rules of specified trigonometric degree.

Definition 1.1. An s -dimensional lattice rule is a cubature formula that can be expressed in the form

$$(1.1) \quad Qf = Q[t, D, Z, s]f \\ := \frac{1}{d_1 d_2 \dots d_t} \sum_{j_1=1}^{d_1} \sum_{j_2=1}^{d_2} \dots \sum_{j_t=1}^{d_t} f \left(\left\{ \frac{j_1 \mathbf{z}_1}{d_1} + \frac{j_2 \mathbf{z}_2}{d_2} + \dots + \frac{j_t \mathbf{z}_t}{d_t} \right\} \right),$$

where d_i are positive integers and $\mathbf{z}_i \in \mathbb{Z}^s$ for all i .

In this theory it is conventional to refer to \mathbb{Z}^s (the set of points all of whose components are integers) as the s -dimensional *unit lattice* denoted by Λ_0^s . The abscissas of the lattice rule Qf lie on an integration lattice Λ , that is, a discrete subset of \mathbb{R}^s that is closed under addition and subtraction and that contains Λ_0^s . The arguments in the right-hand member in (1.1) may be assembled into two matrices.

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These are the $t \times t$ matrix $D = \text{diag}\{d_i\}$ and the $t \times s$ matrix Z whose i th row is \mathbf{z}_i . The rank and invariants of a lattice rule play no major role in the theory treated in this paper, and their definitions are omitted. However, we remark that much of the previous work in this area has been restricted to *rank-1 simple* lattice rules. These are rules that can be expressed in form (1.1) above with $t = 1$ and \mathbf{z}_1 having 1 as its first component.

All cubature rules Q have an abscissa count $N(Q)$ and have a trigonometric degree, say, $d(Q)$. It turns out to be more convenient to work with

$$(1.2) \quad \delta := d + 1,$$

which we term the *enhanced degree*. An *optimal rule* of enhanced degree δ is one whose abscissa count is known to be as small as or smaller than the abscissa count $N(Q')$ of any other rule Q' of this same enhanced degree δ . In this case we denote this count by $N_{\min}(s, \delta)$. A standard goal, which is our ultimate goal, is to find optimal rules.

Optimal rules are already known for $s = 1$ and 2 for all δ ; they are also known for all s with $\delta = 1, 2, 3, 4$ and for $(s, \delta) = (3, 6)$. In each of these cases, at least one of these optimal rules is a lattice rule. Except in the cases just mentioned, no rule is known to be optimal, and it is not known whether there is any case in which one of the optimal rules is not a lattice rule.

A lower bound on $N_{\min}(s, \delta)$, which is based on the character of the set of *moment equations*, appears in [CS96], is denoted here by $N_{ME}(s, \delta)$, and is available for all s and d ; however, except in the aforementioned cases, it is not known whether this bound is attained. In particular:

$$(1.3) \quad \begin{aligned} N_{ME}(1, \delta) &= \delta \\ N_{ME}(2, \delta) &= \delta^2/2 && \delta \text{ even} \\ &= (\delta^2 + 1)/2 && \delta \text{ odd} \\ N_{ME}(3, \delta) &= \delta(\delta^2 + 2)/6 && \delta \text{ even} \\ &= \delta(\delta^2 + 5)/6 && \delta \text{ odd} \\ N_{ME}(4, \delta) &= \delta^2(\delta^2 + 8)/24 && \delta \text{ even} \\ &= (\delta^4 + 14\delta^2 + 9)/24 && \delta \text{ odd} \end{aligned}$$

A completely different bound, valid only for lattice rules, follows from applying Minkowski’s celebrated theorem about admissible lattices to an s -dimensional octahedron $\Omega(s, \delta)$ defined in (2.3). In the present context, this provides a bound $N \geq \delta^s/s!$ for lattice rules. A much deeper result of his introduces a “critical lattice” for $s = 1, 2,$ and 3. The consequence for us is that a bound exists that is specific for lattice rules:

$$(1.4) \quad N \geq \overline{N}_{CL}(s, \delta) := \frac{\delta^s}{s!\theta(s)}.$$

Clearly $\theta(s) \leq 1$. The only known values of $\theta(s)$ are $\theta(1) = \theta(2) = 1$ and $\theta(3) = 18/19$. In the literature on geometry of numbers [GL87], $\theta(s)$ is known as the “density of closest (or densest) lattice packing” for the s -dimensional octahedron. Nontrivial upper bounds for $\theta(s), s \geq 4$, appear to be unknown. Every lattice rule provides a lower bound for $\theta(s)$. Examination of our recent results in Table 2 establishes $\theta(4) \geq \frac{512}{621}$. This improves the result of Klyuchnikov and Reztsov [KR95], $\theta(4) \geq \frac{128}{159}$, by a margin of approximately 0.02. In our context, N is an

integer, so the above inequality may be sharpened to

$$(1.5) \quad N \geq N_{CL}(s, \delta) := \lceil \overline{N}_{CL}(s, \delta) \rceil.$$

For $s = 1$ and 2 , $N_{CL} = N_{ME}$. For $s = 3$ and $\delta = 8$ and ≥ 10 , $N_{CL} \geq N_{ME}$. Numerical values of N_{ME} and N_{CL} are given in Tables 1 and 2.

A small amount of literature (mostly in Russian) has been devoted to *optimal* cubature rules. The optimal rules mentioned above appear in papers by Mysovskikh [Mys85, Mys87, Mys88] and Noskov [Nos85, Nos88a, Nos88b] and are elaborated by Beckers and Cools [BC93], Cools and Sloan [CS96] and Cools and Reztsov [CR97]. The three-dimensional rule is classical and due to Minkowski (see [Fro77] and [Min67, Chapter XIX]).

Furthermore, Noskov and Semenova have published many *nonoptimal* individual three-, four-, and five-dimensional rank-1 simple lattice rules and several families; see, for example, [Nos88a, Nos91, NS96, Sem96]. Each family is a one-parameter system (the parameter being essentially the degree) of rank-1 simple lattice rules; and, since the parameter is unbounded, these include rules of arbitrarily high degree. It is not revealed how they were discovered, but clearly careful effort was expended, and they are far more economical than those (such as the center and vertex rule) previously available. However, it appears that none is likely to be particularly close to optimal. To our knowledge these are the only lattice rules available that are reasonably efficient from the trigonometric point of view.

We have carried out a large-scale computer search with a view to clarifying the situation as far as optimal *lattice* rules in dimensions 3 and 4 are concerned. We have managed to reach degree 30 in three dimensions and to reach degree 24 in four. In this paper, we describe this search and give some background in the context of other analogous searches. We present some of the results.

Our search is however restricted to a subset of the lattice rules, namely, $K(s, \delta)$ of Definition 2.7 below. There are compelling reasons for believing that the optimal lattice rules are members of this set, but this has not been proved. We have come across no counterexample nor any suggestion that such a counterexample may exist. Nevertheless, we retain the distinction and refer to the optimal lattice rules of this set as K -optimal lattice rules.

2. UNDERLYING THEORY

The theory on which our search is based is closely analogous to the theory on which some searches for good lattices are based. We give a brief description here, mainly to introduce the standard notation.

A lattice Λ may be defined in terms of an $s \times s$ matrix A known as a *generator matrix*. This means that all elements of Λ are of the form $x = \lambda A$, where $\lambda \in \mathbb{Z}^s$. The *dual* lattice Λ^\perp may be defined as one having generator matrix $B = (A^T)^{-1}$. The reader will recall that, since Λ is an integration lattice, that is, $\Lambda \supseteq \Lambda_0^s$, its dual Λ^\perp is an integer lattice and may be generated by an integer-valued matrix B .

When U is any unimodular matrix, $H = UB$ is also a generator matrix for Λ^\perp . For any given B , there exists a particular choice for U that will provide a generator matrix $H = UB$ that is in Hermite normal form (utlf). That is,

$$(2.1) \quad \begin{aligned} H_{c,c} &> 0 \\ H_{r,c} &= 0 && r > c \\ H_{r,c} &\in [0, H_{c,c}) && r < c. \end{aligned}$$

A (1-1) correspondence exists between the set of s -dimensional lattice rules and the set of $s \times s$ matrices in Hermite normal form. This has been exploited previously to organize searches [LS93] but is not exploited in that way here.

The quantity $|\det B|$ is conventionally known as the *order* of the lattice Λ^\perp . The simplex whose $s + 1$ vertices make up the s rows of B , namely, \mathbf{b}_j ($j = 1, 2, \dots, s$), together with the origin \mathbf{O} is known as a *basic cell* of Λ^\perp . (Any simplex obtained in this way using any generating matrix UB is also a basic cell, as is any simplex obtained by translating one of these simplices.) The s -volume of a basic cell is $|\det B|/s!$. In fact, all s -dimensional simplectical regions whose vertices are distinct elements of Λ^\perp have s -volume $k|\det B|/s!$, where k is some nonnegative integer. (Any set of vertices for which $k = 1$ forms a basic cell.) The relevance of the basic cell to our search lies in the fact that the abscissa count of Q coincides with $|\det B|$ (see [Lyn89]), that is,

$$N(Q(\Lambda)) = |\det B| = \prod_{i=1}^s H_{i,i}.$$

This may be reexpressed as follows.

Theorem 2.1. *The abscissa count N of $Q(\Lambda)$ coincides with the order of Λ^\perp .*

When $Q(\Lambda)$ is the lattice rule whose integration lattice is Λ , the associated Poisson summation formula reduces to an expression for the discretization error, namely,

$$E_{Q(\Lambda)}f := Q(\Lambda)f - If = \sum_{\substack{\mathbf{h} \in \Lambda^\perp \\ \mathbf{h} \neq \mathbf{0}}} \hat{f}_{\mathbf{h}},$$

where $\hat{f}_{\mathbf{h}}$ is the Fourier coefficient of f and Λ^\perp is the dual lattice of Λ . When f is a trigonometric polynomial of degree d or less, $\hat{f}_{\mathbf{h}} = 0$ when $|\mathbf{h}| > d$, so all but a finite set of terms in this sum vanish. Thus, the condition that $Q(\Lambda)f$ is exact for these polynomials reduces to the condition that Λ^\perp has no elements, other than the origin itself, in the region $|\mathbf{h}| \leq d$, which we denote by $\Omega(s, d)$. We may restate this as follows:

$$(2.2) \quad \delta(Q(\Lambda)) := d(Q(\Lambda)) + 1 = \min_{\substack{\mathbf{h} \in \Lambda^\perp \\ \mathbf{h} \neq \mathbf{0}}} |\mathbf{h}|.$$

This equation relates the location of points $\mathbf{h} \in \Lambda^\perp$ with the enhanced degree δ of $Q(\Lambda)$. We may use classical terminology to reexpress the import of this equation in terms taken from the geometry of numbers [GL87].

Definition 2.2. (Classical) A lattice L is “admissible” with respect to a region Ω if all its elements (other than the origin) lie outside Ω .

Such a lattice is conventionally known as an Ω -admissible lattice. Applied to our region

$$(2.3) \quad \mathbf{h} \in \Omega(s, \delta) \text{ when } |\mathbf{h}| \leq \delta,$$

we have the following definition.

Definition 2.3. An $\Omega(s, \delta)$ -admissible lattice is an integer lattice having no elements, other than the origin, in the interior of $\Omega(s, \delta)$.

Using this terminology, we may write the content of (2.2) as follows:

Theorem 2.4. $Q(\Lambda)$ is of enhanced degree δ or greater if and only if Λ^\perp is $\Omega(s, \delta)$ -admissible.

This theorem, together with Theorem 2.1, leads to the following geometric characterization.

Theorem 2.5. $Q(\Lambda)$ is an optimal lattice rule of enhanced degree δ when Λ^\perp is an $\Omega(s, \delta)$ -admissible lattice and no other $\Omega(s, \delta)$ -admissible lattice has a lower order.

In passing, it is pertinent to mention that many other criteria are in use to characterise efficient cubature rules, and that some, like the enhanced degree in (1.2) above, are based on exact evaluation of specified sets of Fourier coefficients. Some of these latter are discussed in Lyness [Lyn88] and may be described in terms of Ω -admissible lattices with Ω redefined appropriately. The two most familiar choices are illustrated in, e.g., [BC93]. Other choices are investigated in [CR97] and [LS97].

We now return to the problem at hand. In this paper, Ω is defined in (2.3) and we are treating the enhanced degree, defined in (1.2).

A dynamic approach to the problem of finding an optimal rule might involve perturbing any given $\Omega(s, \delta)$ -admissible lattice Λ^\perp , with a view to reducing the s -volume of its unit cell but keeping it $\Omega(s, \delta)$ -admissible, that is, not allowing any lattice point to enter the fixed region $\Omega(s, \delta)$.

It is reasonable to believe that the process of making this unit cell small, that is, making the lattice Λ^\perp denser and reducing its order, would, in general, move lattice points towards the origin. This process would be seriously inhibited by the boundary of $\Omega(s, \delta)$. Ultimately, (as the wiggle room disappears) one would expect progress to come to a complete stop (grind to a halt) at a stage where many points of Λ^\perp were (jammed) on this boundary. Thus, it is plausible to believe that the lattice Λ of an optimal lattice rule $Q(\Lambda)$ of enhanced degree δ will have a dual lattice Λ^\perp with many elements on this boundary. The underlying feature of our search is that it is limited to dual lattices having this property.

The $(s - 1)$ -dimensional *facet-pair* of an s -crosspolytope is the s -dimensional generalization of a two-dimensional pair of opposite faces of a regular (three-dimensional) octahedron. We recall the following notation:

$$|\mathbf{x}| = |(x_1, x_2, x_3, \dots, x_s)| = |x_1| + |x_2| + |x_3| + \dots + |x_s|$$

$$\mathbf{h} \in \Omega(s, \delta) \text{ when } |\mathbf{h}| \leq \delta$$

$$\mathbf{h} \in \bar{\Omega}(s, \delta) \text{ when } |\mathbf{h}| = \delta.$$

In the sequel, σ_i stands for $+1$ or for -1 .

Definition 2.6. The facet-pair $F(\delta, \sigma_1, \sigma_2, \sigma_3, \dots, \sigma_s)$ comprises \mathbf{h} satisfying

$$\mathbf{h} \in \bar{\Omega}(s, \delta) \text{ and}$$

either

$$h_i = \sigma_i |h_i| \text{ for all } i = 1, 2, 3, \dots, s$$

or

$$h_i = -\sigma_i |h_i| \text{ for all } i = 1, 2, 3, \dots, s.$$

Definition 2.7. The population $K(s, \delta)$ comprises all s -dimensional lattices that may be *generated* by s point pairs, each of which belongs to a distinct $(s - 1)$ -dimensional facet-pair of the s -octahedron (s -crosspolytope) $\Omega(s, \delta)$.

Note that a lattice in $K(s, \delta)$ cannot have enhanced degree exceeding δ because, by definition, it includes points \mathbf{h} having $|\mathbf{h}| = \delta$. In general such a lattice is of degree less than δ .

We search this population for the rule or rules defined as follows.

Definition 2.8. A $K(s, \delta)$ -optimal rule is a rule of minimum abscissa count among those of enhanced degree δ whose dual lattice Λ^\perp is in $K(s, \delta)$.

In the next section, we shall require subsets of $K(s, \delta)$. These will be denoted by $K(s, \delta; X)$, where X will identify the particular subset in question.

3. THE SEARCH PROGRAMS

In this section we describe the implementation of programs based on the ideas and definitions introduced at the end of the preceding section. It has turned out that the four-dimensional program is significantly more complicated than the three-dimensional program. For this reason, after introducing some common s -dimensional notation, we describe the three-dimensional program first. Then, with the underlying ideas exposed in the simpler context, we treat the four-dimensional program.

In three or more dimensions, significant effort can be saved by exploiting the existence of sets of symmetrically equivalent lattices. A group of linear transformations takes the s -cube, or the s -octahedron, into itself. Applying one of these transformations to a rule or a lattice provides another (generally different) rule or lattice having the same geometric characteristics. Naturally, two lattices related in this way have the same (enhanced) degree and the same order (abscissa count). A set of symmetrically equivalent lattices may have as many as $s!2^{s-1}$ members. Once one member of such a set is established to be optimal, the other members of the set may be rapidly identified and are also optimal. Thus, if we are able to subdivide the search population in such a way that a search over one part will recover only symmetric equivalents of a search over another part, we may exploit this by searching only one of these parts. A search over the second part can be safely omitted, as it would reveal only optimal lattices that are symmetric equivalents of optimal lattices already identified.

In three dimensions, it is particularly easy to exploit the concept of sets of symmetrically equivalent lattices. In view of Definition 2.7 above, the set $K(3, \delta)$ includes all lattices generated by three points $\mathbf{b}_1, \mathbf{b}_2,$ and \mathbf{b}_3 , where each lies on a different facet-pair. We define a subset of $K(3, \delta)$, which we denote by K^* . This includes only lattices generated by

$$(3.1) \quad \begin{aligned} \mathbf{b}_1 &\in F(\delta, +, +, +) \\ \mathbf{b}_2 &\in F(\delta, +, +, -) \\ \mathbf{b}_3 &\in F(\delta, +, -, +). \end{aligned}$$

It is straightforward to show that all lattices in $K(3, \delta)$ have a symmetrically equivalent lattice in K^* . Thus, we may restrict our search to the elements of K^* and then include, in addition, all symmetric equivalents. The outcome is the same as

if we had treated all the elements of $K(3, \delta)$, but is obtained at approximately one fourth the cost. (The corresponding statement in four dimensions is not true.)

Our *search module* has two principal modes of operation. In mode 1 (its usual mode) it requires as input numerical values of δ and N_L and N_U . It also requires a specification of the population to be treated. (When $s = 3$, this is simply the set K^* discussed above. For $s = 4$, as described later, several different population specifications may be used in different runs.) It carries out a search over this population set and either

- (A) provides the generator matrix of a lattice Λ^\perp for which the rule $Q(\Lambda)$ is of enhanced degree δ , the order N of this rule satisfies $N \in [N_L, N_U]$, and there is no rule of lower order in this interval; or
- (B) reports that no lattice Λ^\perp of enhanced degree δ with $N \in [N_L, N_U]$ exists in the specified input population.

To obtain this information, the search module proceeds as follows. It carries out a loop over all matrices B whose rows $\mathbf{b}_1, \mathbf{b}_2, \dots, \mathbf{b}_s$ are elements of their respective facet-pairs (see (3.1) above). Thus, there are possibly $\binom{s + \delta - 1}{s - 1}^s$ matrices B to consider. For each, the order $|\det B|$ is evaluated. Unless $|\det B| \in [N_L, N_U]$, this matrix B is abandoned, and the next matrix B is treated.

In the relatively few cases in which $|\det B|$ is within these limits, an algorithm for determining the enhanced degree of Λ^\perp (or an upper bound on this) is invoked. Unless this enhanced degree is δ , this matrix B is abandoned and the next one is treated. Should this enhanced degree turn out to be δ , ipso facto one lattice satisfying (A) above is available. In mode 1, the search immediately downgrades N_U to $N - 1$ and continues (unless $N = N_L$, in which case it stops).

In all cases, if the module encounters no $\Omega(s, \delta)$ -admissible lattice of enhanced degree δ , the conclusion (B) above is reported.

The module can also be run in mode 2. This requires the same input as in mode 1. However, instead of downgrading N_U to $N - 1$ when one lattice satisfying (A) is encountered, it downgrades N_U to N and continues until all matrices B have been treated. This mode is normally used when the optimal N_{opt} has already been determined and is invoked to see whether there are several different solutions. One sets $N_L = N_U = N_{opt}$.

The list of rules in Table 3 was obtained as follows. For each value of δ , the search module was used with N_U large and $N_L = \max(N_{ME}(3, \delta), N_{CL}(3, \delta))$ as given in (1.3) and (1.5). The value of N returned in item (A) was used in a second run using mode 2. Finally, the list of matrices was processed to remove all symmetric equivalents. Note that, without the second run, one of the entries for each of $\delta = 5$ and 11 in Table 3 would have been missed.

The 4-octahedron has eight facet-pairs.

F_0	$F(\delta, +, +, +, +)$	E
F_1	$F(\delta, -, +, +, +)$	O
F_2	$F(\delta, +, -, +, +)$	O
F_3	$F(\delta, -, -, +, +)$	E
F_4	$F(\delta, +, +, -, +)$	O
F_5	$F(\delta, -, +, -, +)$	E
F_6	$F(\delta, +, -, -, +)$	E
F_7	$F(\delta, -, -, -, +)$	O

Each has been assigned a serial number, which appears as a subscript in column 1. For later convenience, in column 3 we have assigned a parity to each. $F(\delta, \sigma_1, \sigma_2, \sigma_3, \sigma_4)$ is of even parity E if the set $(\sigma_1, \sigma_2, \sigma_3, \sigma_4)$ contains an even number of elements +1.

Every lattice Λ^\perp in $K(4, \delta)$ is generated by four points on four distinct facet-pairs. We term such a set of facet-pairs a *quartet (of facet-pairs)*. If we were to take no account of the symmetric equivalents, we would need to treat every distinct quartet separately. There are seventy distinct quartets, this being the number of ways of choosing four facet-pairs from the total of eight facet-pairs listed above. The following discussion is devoted solely to establishing Theorem 3.4 below, which assures us that only four of these quartets need be searched to ensure that we recover at least one symmetric equivalent of every optimal rule.

Definition 3.1. An individual quartet, denoted by $q(N_1, N_2, N_3, N_4)$ where $0 \leq N_1 < N_2 < N_3 < N_4 \leq 7$ comprises a set of four distinct facet-pairs $F_{N_1}, F_{N_2}, F_{N_3}$ and F_{N_4} .

The *type* of a quartet q is $\min(NE, NO)$ where NE is the number of even facet-pairs and NO is the number of odd facet-pairs in q . For example, $q(0, 4, 5, 6)$ contains three even facet-pairs, namely, $F_0, F_5,$ and F_6 , together with one odd facet-pair, F_4 . Thus its type is 1, this being the minimum of $NE = 3$ and $NO = 1$.

Definition 3.2. Let $q(N_1, N_2, N_3, N_4)$ be one of these 70 quartets. The population $K(4, \delta; q)$ comprises any lattice that may be generated by four points \mathbf{b}_i , where $\mathbf{b}_i \in F_{N_i}, i = 1, 2, 3, 4$.

The union of all seventy of these populations $K(4, \delta; q)$ includes all lattices that may be generated by four distinct points, each of which lies on a distinct facet-pair of the 4-octahedron, and so coincides with $K(4, \delta)$.

Let \mathcal{G}_i be an element of the group \mathcal{G} of 384 affine transformations that take the 4-octahedron into itself. Specifically, this transformation takes any facet-pair F_{N_j} into some other facet-pair F_{N_k} , which we may denote by $\mathcal{G}_i F_{N_j}$. By the same token, this transformation takes separately each of a set of four facet-pairs into another set of four facet-pairs.

Definition 3.3. Let $q = q(N_1, N_2, N_3, N_4)$ be one of these 70 quartets. The quartet comprising the four facet-pairs $\mathcal{G}_i F_{N_j}, j = 1, 2, 3, 4$, is termed a symmetric copy of $q(N_1, N_2, N_3, N_4)$ and is denoted by $\mathcal{G}_i q$.

Let q_1 stand for the quartet $q(0, 2, 4, 7)$. Clearly, a search over $K(4, \delta; \mathcal{G}_i q_1)$ will yield only lattices that are symmetrically equivalent to those obtained in the same search over $K(4, \delta; q_1)$. It is a trivial calculation to obtain all symmetric equivalents of a particular lattice. Thus, carrying out a search over more than one quartet belonging to the set of quartets $\mathcal{G}_i q_1$ is unnecessary. A straightforward calculation (elaborated in the Appendix) reveals that there are only 32 distinct quartets of this form. Thus, S_1 is a set of order 32, and we need to search over only one of these 32 quartets. Our choice for q_1 could be replaced by any other member of S_1 with the same result.

We repeat this operation starting with the three specific quartets given in the theorem.

Theorem 3.4. Let $q_0 = q(1, 2, 4, 7)$, $q_1 = q(0, 2, 4, 7)$, $q_{2a} = q(0, 2, 4, 6)$, and $q_{2b} = q(0, 2, 4, 5)$ and the sets of quartets constituting symmetric copies of q_j be denoted

by S_j . Then the sets S_i are mutually disjoint. They are of orders 2, 32, 12, and 24, respectively, and their union includes all seventy quartets.

Proof. The sets S_0, S_1, S_{2a}, S_{2b} , are listed in the Appendix. The theorem may be verified by four sets of 384 simple calculations. One calculates $q = \mathcal{G}_i q_j$ ($i = 1, \dots, 384$), $j = 0, 1, 2a, 2b$, and verifies that each is a member of the expected set S_j . The reader will note that all elements of $S_i, i = 0, 1$, are of type i and elements of S_{2a} and S_{2b} are of type 2. It is straightforward to show that none of the 384 transformations alters the type of the quartet. The Appendix provides further details. \square

The results of our computer searches for four-dimensional optimal rules are presented in Table 5. This is in three parts. For each value of $\delta \in [1, 13]$ we have made four distinct runs and (unless there are calculational errors) we have a complete list of all optimal $K(4, \delta)$ rules.

For $\delta \in [14, 17]$ we reduced the population to $K(4, \delta; q_1)$, where, as before, $q_1 = q(0, 2, 4, 7)$. This restriction to a single quartet reduces the overall run time by a factor of 4.

Beyond $\delta = 18$, even this became too time consuming, and we reduced the population once more to $K(4, \delta; q_1^+)$. The symbol q_1^+ is used here to denote a subset of q_1 that includes all of F_2, F_4 , and F_7 , but only the part of $F_0 = F(\delta, +, +, +, +)$ for which $x_1 \geq x_2 \geq x_3 \geq x_4$. This reduces the size of the population by a factor of up to 24. But almost certainly some optimal rules are missed.

We have described the three searches above in terms of the results. In the order of implementation, we first carried out a search using population $K(4, \delta; q_1^+)$ for δ up to 24. Next, we used $K(4, \delta; q_1)$ for δ up to 17. Finally, we carried out a complete search, using four choices for q , for δ up to 13.

4. NEW RESULTS

In subsections 4.2 and 4.3, we present some of our three- and four-dimensional results, respectively. Subsection 4.1 is devoted to careful definitions of the notation used in the tables.

4.1. Abscissa counts. In this first subsection we present the progress toward determining $N_{opt}(s, \delta)$, the optimal abscissa count for any s -dimensional rule of enhanced trigonometric degree δ . We have in general obtained well-defined bounds on this quantity. These are denoted by $N_X(s, \delta)$, where the subscript X indicates a limitation to the class of rules considered.

The five principal abscissa count functions we have listed are as follows:

- N_{ME} : A theoretical lower bound for any rule of enhanced degree δ , based on the relevant *Moment Equations*.
- N_{CL} : The Minkowski lower bound for any lattice rule of enhanced degree δ , based on the existence of the *critical lattice* (known only for dimensions $s = 1, 2$ and 3).
- N_{KO} : The lowest count for any $K(s, \delta)$ -optimal rule. (We also list variants of N_{KO} .)
- N_{r1s} : The lowest abscissa count for any optimal rank-1 simple rule.
- N_{prev} : The lowest abscissa count for any rule published in references [Nos88a, NS96]. These are all rank-1 simple.

TABLE 1. Three-dimensional abscissa counts

δ $= d + 1$	N_{ME}	N_{CL}	N_{KO}	N_{r1s}	N_{prev}	$\rho(N_{KO})$
1	1	1	1	1	1	0.167
2	2	2	2	2	2	0.667
3	7	5	7	7	7	0.643
4	12	12	12	12	12	0.889
5	25	22	27	27	27	0.772
6	38	38	38	38	38	<u>0.947</u>
7	63	61	70	70	70	0.817
8	88	91	92	92	92	0.928
9	129	129	144	145	145	0.844
10	170	176	178	178	178	0.936
11	231	235	260	260	260	0.853
12	292	304	304	312	312	<u>0.947</u>
13	377	387	421	421	421	0.870
14	462	483	486	486	486	0.941
15	575	594	635	635	635	0.886
16	688	721	724	724	724	0.943
17	833	865	921	921	921	0.889
18	978	1026	1026	1038	1038	<u>0.947</u>
19	1159	1207	1276	1276	1319	0.896
20	1340	1408	1412	1412	1412	0.944
21	1561	1630	1708	1723	1771	0.904
22	1782	1874	1878	1878	1942	0.945
23	2047	2141	2240	2255	2327	0.905
24	2312	2432	2432	2448	2532	<u>0.947</u>
25	2625	2749	2865	2865	2977	0.909
26	2938	3093	3098	3098	3218	0.946
27	3303	3463	3591	3591	3751	0.914
28	3668	3862	3868	3868	4032	0.946
29	4089	4291	4445	4445	4635	0.915
30	4510	4750	4750	4770	4958	<u>0.947</u>

Formulas for N_{ME} are given for all (s, δ) in reference [CS96] and repeated by us for $s \leq 4$ in (1.3) above. N_{CL} is simply (1.5) above. The principal contribution of our work is the list of values of N_{KO} and some variants in Tables 1 and 2. We obtained the fourth abscissa count N_{r1s} for $s = 3$ ($\delta \leq 30$) and $s = 4$ ($\delta \leq 13$) using a simple search program not discussed here. The fifth abscissa count N_{prev} is readily gleaned from the cited literature.

The three-dimensional abscissa counts listed in Table 1 are all precisely as defined above. The four-dimensional abscissa counts listed in Table 2 are also precisely as defined above for $\delta \leq 13$. For higher values of δ , the entries under N_{KO} refer to the results of restricted searches, as indicated in Table 5 and specified at the end of the preceding section. The corresponding entries under N_{r1s} may not be optimal. Rules corresponding to every abscissa count given in the columns labeled N_{KO} and N_{r1s} are specified in Tables 3, 4, 5, and 6.

For odd $\delta > 14$ some rules have been published, but these use more points than published rules of higher degree. We have omitted these.

TABLE 2. Four-dimensional abscissa counts

δ	N_{ME}	N_{KO}	N_{r1s}	N_{prev}	$\rho(N_{KO})$
1	1		1	1	0.042
2	2		2	2	0.333
3	9		9	9	0.375
4	16		16	16	0.667
5	41	45	46	46	0.579
6	66	68	70	70	0.794
7	129	152	152	156	0.658
8	192	212	212	212	0.805
9	321	375	398	414	0.729
10	450	516	522	522	0.807
11	681	857	857	1076	0.712
12	912	1064	1092	1092	0.812
13	1289	1601	1601	1709	0.743
14	1666	1958	[1958]	3075	0.818
15	2241	2834	[2834]		0.744
16	2816	3312	[3376]	3522	<u>0.824</u>
17	3649	4628	[4633]		0.752
18	4482	5354	[5354]	6242	0.817
19	5641	7081	[7081]		0.767
20	6800	8148	[8148]	8840	0.818
21	8361	10552	[10552]		0.768
22	9922	11886	[11886]	14102	0.821
23	11969	15154	[15154]		0.769
24	14016	16812	[17208]		0.822

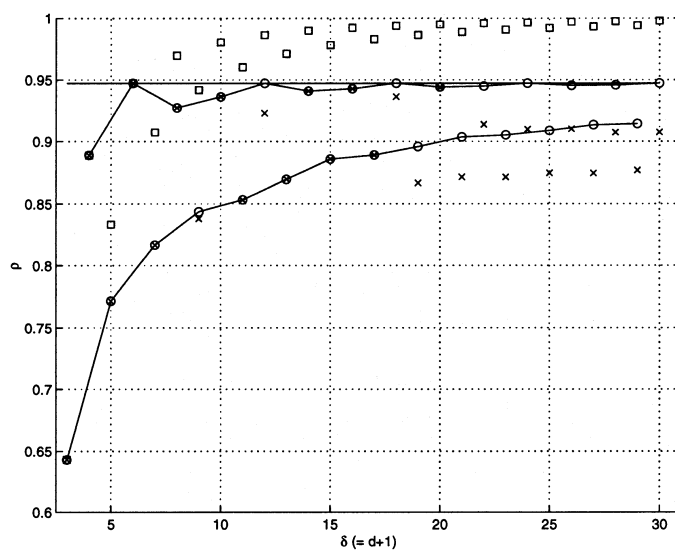


FIGURE 1. ρ as a function of δ for three-dimensional rules (\times refers to N_{prev} , \circ refers to N_{KO} , \square refers to N_{ME} , the line at $\rho = 18/19$ refers to N_{CL})

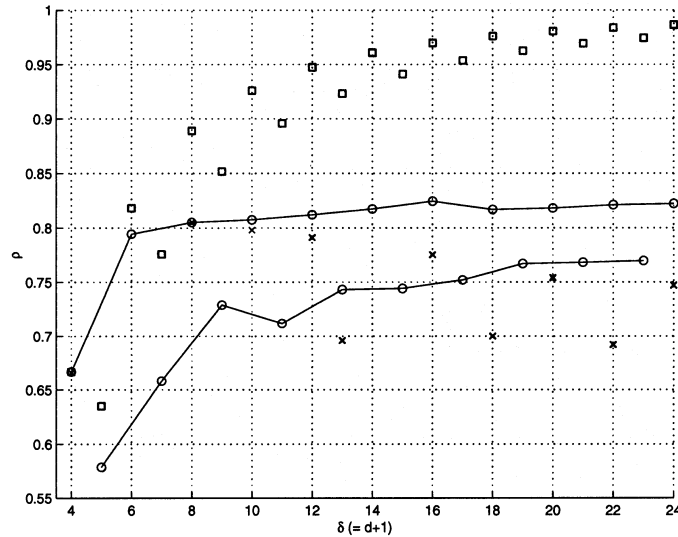


FIGURE 2. ρ as a function of δ for four-dimensional rules (\times refers to N_{prev} , \circ refers to N_{KO} , \square refers to N_{ME})

In Figures 1 and 2 we present much of the material in Tables 1 and 2 graphically. For any abscissa count N , we can calculate the associated *packing factor*

$$(4.1) \quad \rho(N) := \frac{\delta^s}{s!N}.$$

This is a measure of the efficiency of any rule $Q(\Lambda)$ of enhanced degree δ and abscissa count N and is the packing factor of the dual lattice Λ^\perp . The packing factor is bounded by $\theta(s)$. In the final section we shall illustrate our discussion of some of these results using these figures.

Many of the entries in the tables specify rank-1 simple rules. When $Q(\Lambda)$ is an s -dimensional rank-1 simple rule, the Hermite normal form (see (2.1) above) of the generator matrix of Λ^\perp has a readily recognisable form as its principal minor coincides with the identity matrix. The $D - Z$ form (see (1.1)) of this rule is then $Q[1, D, \mathbf{z}, s]$ with

$$D = N = H_{s,s}; \quad \mathbf{z} = (N - H_{1,s}, N - H_{2,s}, \dots, N - H_{(s-1),s}, 1).$$

Naturally, this is in the same equivalence class as the rule specified by

$$D = N = H_{s,s}; \quad \mathbf{z} = (1, H_{1,s}, H_{2,s}, \dots, H_{(s-1),s}).$$

4.2. Three-dimensional lattice rules. For every abscissa count we have listed, we have specified at least one cubature rule. Table 3 contains specifications of thirty-one K -optimal rules. This list is complete in the sense that every K -optimal rule of enhanced degree thirty or less is included here or is symmetrically equivalent to one listed here. This specification comprises the nontrivial elements of the Hermite normal form of Λ^\perp (unique to the rule). See (2.1) above.

TABLE 3. Three-dimensional K -optimal lattice rules

δ	N	Hermite Normal Form of Dual Lattice						μ	Rank
		H_{11}	H_{12}	H_{13}	H_{22}	H_{23}	H_{33}		
2	2	1	0	1	1	1	2	1	1
3	7	1	0	2	1	3	7	8	1
4	12	1	0	3	1	5	12	12	1
5	27	1	0	4	1	10	27	24	1
		1	1	4	3	6	9	4	2
6	38	1	0	7	1	11	38	8	1
7	70	1	0	16	1	25	70	24	1
8	92	1	0	9	1	39	92	24	1
9	144	1	1	11	4	16	36	8	2
10	178	1	0	11	1	75	178	24	1
11	260	1	0	40	1	94	260	24	1
		1	0	48	2	56	130	24	2
12	304	2	0	14	2	22	76	8	3
13	421	1	0	16	1	182	421	24	1
14	486	1	0	41	1	57	486	24	1
15	635	1	0	146	1	274	635	8	1
16	724	1	0	49	1	79	724	24	1
17	921	1	0	81	1	222	921	24	1
18	1026	3	0	21	3	33	114	8	3
19	1276	1	0	222	1	538	1276	24	1
20	1412	1	0	59	1	665	1412	24	1
21	1708	1	1	121	2	338	854	8	2
22	1878	1	0	75	1	731	1878	24	1
23	2240	1	0	166	4	255	560	24	1 (not simple)
24	2432	4	0	28	4	44	152	8	3
25	2865	1	0	222	1	965	2865	24	1
26	3098	1	0	423	1	1299	3098	24	1
27	3591	1	0	278	1	1718	3591	8	1
28	3868	1	0	205	1	975	3868	24	1
29	4445	1	0	750	1	1635	4445	24	1
30	4750	5	0	35	5	55	190	8	3

The penultimate column contains μ , the number of distinct rules (symmetric copies) in the symmetry group that contains the listed rule. These may be obtained from the listed rule by coordinate reversal and interchange. Naturally, we list only one rule of the μ possibilities. This is chosen to be the first in a lexicographic ordering based on the diagonal elements, followed by the nondiagonal elements in the order used in the table. In the language of [LS93], this provides a *senior*. Also, if the rank is 1, this provides a rank-1 *simple* rule, unless there happens to be no rank-1 simple rule in the set.

The eight rules of enhanced degree $\delta = 6k$ with $k > 1$ are simply k -copy versions of the eight rules of enhanced degree 6. These are of rank 3.

A supplementary list of three-dimensional optimal rank-1 simple lattice rules is given in Table 4. This list is of the same character as the previous list. It includes all optimal rank-1 simple rules for those degrees for which such a rule does not appear in the previous list.

TABLE 4. Three-dimensional optimal rank-1 simple lattice rules

δ	N	Hermite Normal Form of Dual Lattice						μ
		H_{11}	H_{12}	H_{13}	H_{22}	H_{23}	H_{33}	
9	145	1	0	9	1	61	145	24
12	312	1	0	13	1	115	312	24
		1	0	29	1	67	312	24
18	1038	1	0	35	1	365	1038	24
		1	0	119	1	421	1038	24
21	1723	1	0	24	1	464	1723	24
		1	0	79	1	755	1723	24
23	2255	1	0	100	1	172	2255	24
24	2448	1	0	185	1	1081	2448	24
		1	0	199	1	479	2448	24
30	4770	1	0	131	1	689	4770	24
		1	0	101	1	1339	4770	24

4.3. **Four-dimensional lattice rules.** We have reported our four-dimensional results in almost the same way as the three-dimensional results. The differences arise from having to curtail our effort because of the higher computational expense. As in the three-dimensional case, we have specified in Table 5 the optimal rules we have found. As mentioned in Section 3, any of these may be actual optimal rules of the stated enhanced degree. We have found all the K -optimal rules for $\delta \in [1, 13]$, all the $K(4, \delta; q_1)$ -optimal rules for $\delta \in [14, 17]$, and all the $K(4, \delta; q_1^+)$ -optimal rules

TABLE 5. Four-dimensional K -optimal lattice rules

δ	N	Hermite Normal Form of Dual Lattice										μ	Rank
		H_{11}	H_{12}	H_{13}	H_{14}	H_{22}	H_{23}	H_{24}	H_{33}	H_{34}	H_{44}		
Full Search over $K(4, \delta)$													
1	1	1	0	0	0	1	0	0	1	0	1	1	1
2	2	1	0	0	1	1	0	1	1	1	1	2	1
3	9	1	0	1	1	1	1	2	3	0	3	8	2
		1	0	0	2	1	0	3	1	4	9	64	1
4	16	1	1	1	1	2	0	2	2	2	4	2	3
		1	0	1	2	1	2	1	4	0	4	12	2
		1	0	0	3	1	1	2	2	6	8	24	2
		1	0	0	3	1	0	5	1	7	16	48	1
5	45	1	0	0	4	1	1	6	3	9	15	24	2
6	68	1	0	0	13	1	1	6	2	16	34	48	2
7	152	1	0	0	16	1	0	28	1	37	152	96	1
8	212	1	0	0	9	1	0	33	1	87	212	192	1
9	375	1	1	1	6	5	0	10	5	10	15	24	3
10	516	1	0	0	15	1	0	83	2	118	258	192	2
11	857	1	0	0	188	1	0	207	1	351	857	48	1
12	1064	1	0	0	153	1	0	259	2	98	532	96	2
13	1601	1	0	0	40	1	0	310	1	408	1601	48	1
Full Search over $K(4, \delta; q_1)$													
14	1958	1	0	0	107	1	0	229	1	525	1958	192	1
15	2834	1	0	0	892	1	0	1123	1	1314	2834	96	1
		1	0	0	294	1	1	117	2	507	1417	96	1
16	3312	1	0	0	495	1	0	737	2	450	1656	96	2
17	4628	1	0	0	1123	1	1	327	2	1032	2314	96	2
Full Search over $K(4, \delta; q_1^+)$													
18	5354	1	0	0	83	1	0	1253	1	1863	5354	192	1
19	7081	1	0	0	241	1	0	1433	1	1616	7081	48	1
20	8148	1	0	0	371	1	0	1401	1	3299	8148	192	1
21	10552	1	0	0	1670	1	0	2111	1	2746	10552	192	1
22	11886	1	0	0	457	1	0	3753	1	4079	11886	192	1
23	15154	1	0	0	2602	1	0	6037	1	6424	15154	96	1
24	16812	1	0	0	109	1	1	1717	3	1677	5604	192	2

TABLE 6. Four-dimensional rank-1 simple lattice rules

δ	N	Hermite Normal Form of Dual Lattice										μ
		H_{11}	H_{12}	H_{13}	H_{14}	H_{22}	H_{23}	H_{24}	H_{33}	H_{34}	H_{44}	
9	398	1	0	0	8	1	0	61	1	149	398	192
16	3376	1	0	0	169	1	0	1091	1	1387	3376	192
17	4633	1	0	0	547	1	0	1936	1	1965	4633	48
24	17208	1	0	0	919	1	0	4701	1	5557	17208	192

for $\delta \in [18, 24]$. The supplementary Table 6 simply specifies some rank-1 simple rules whose abscissa counts appear in Table 2 but are not specified elsewhere.

5. FURTHER COMMENTS

Any historical perspective on rules of specified trigonometrical degree would mention the widespread use of the product trapezoidal rule, and the center and vertex rule since the beginning of the twentieth century. However, the serious study of such rules seems to have started in the final fifteen years of that century. The earlier work of this period, mainly by Russian authors, has been strictly limited to rank-1 simple rules. They have produced and established the optimal degree rules up to $\delta = 4$. These authors have been concerned mainly with rule families in three, four, and five dimensions. Each family contains rules of arbitrarily high degree. Other economical rules seem to have been provided only as spin-off, and no claim has been made for optimality. However, in retrospect we have ascertained that in three dimensions their rules are optimal rank-1 simple rules for all $\delta \leq 18$ but that in four dimensions, they are optimal only for odd δ up to 5 and for even δ up to 12.

To our knowledge, the only other set of rules proposed in this context are the Smolyak rules [CNR99]. These were designed for high dimensions and high degrees. In three and four dimensions and for values of δ considered here, the K -optimal rules presented here are well over ten times more cost effective than the corresponding Smolyak rules.

Figures 1 and 2 illustrate most of the abscissa counts listed in Tables 1 and 2. We note the dichotomy between even and odd degree, which seems to occur in both the theoretical limit N_{ME} and results such as N_{KO} and N_{r1s} (not shown in figures but reported in the tables) and N_{prev} .

As discussed in Section 1, we have no theory to exclude the possibility that, for larger δ , the optimal rule of trigonometric degree δ is not a lattice rule. If this were the case, in Figures 1 and 2 there would be missing entries above the lines joining the circles, but below the theoretical limit represented by squares. Also unsatisfactory is the fact that we cannot establish that the K -optimal lattice rule is actually an optimal lattice rule. This is more frustrating because the anecdotal evidence is overwhelming. We have several incomplete proofs, characterized by our inability to bridge in each case what seems to be a minor lacuna. However, we have an example of a rule that is $K(4, \delta)$ -optimal, but not $K(4, \delta; q_2)$ -optimal. The 375 point lattice listed in Table 5 for $\delta = 9$ is not in $K(4, 9; q_2)$. The $K(4, 9; q_2)$ -optimal rules have an abscissa count of 390. And we have encountered many examples in which the restriction to $K(4, \delta; q_1^+)$ has resulted in missing some excellent rules.

One of the unsatisfactory features of our approach is its high computational cost. We have derived a somewhat unrealistic upper bound on the complexity. This depends in the first place on ν , the number of distinct generator matrices we start with. As specified in Section 3,

$$\nu = \binom{s + \delta - 1}{s - 1}^s = \mathcal{O}(\delta^{s^2 - s})$$

for fixed s and increasing δ .

Only a proportion that appears to decrease with increasing δ is treated further to find N . After this, a minute proportion of these are retained to find their degree. A simple basic form of our algorithm to determine the degree of a lattice rule requires time proportional to $\bar{\delta}^{s-1}$, where $\bar{\delta}$ is the degree of the lattice. In fact, all but a handful have degree strictly less than $\bar{\delta}$. To obtain a complexity bound, we replace both proportions by 1 and replace $\bar{\delta}$ by δ . This approach leads to a complexity bounded above by δ^{s^2-1} .

For the values of δ for which we carried out careful timing checks, the computational cost does increase very rapidly with increasing δ , although not nearly so rapidly as the complexity bound derived above might suggest. To give the reader an idea, we list some timings below for a particular processor.¹

- For $s = 3$ all $\delta \leq 30$ are treated within 33 minutes.
- For $s = 4$ all $\delta \leq 8$ are treated within 34 minutes.
- For $s = 4$ and $\delta = 10$ the search required 6.5 hours.
- For $s = 4$ and $\delta = 14$ the search restricted to $K(4, 14; q_1)$ required 120 hours.
- For $s = 4$ and $\delta = 17$ the search restricted to $K(4, 17; q_1^+)$ required 145 hours.
- For $s = 4$ and $\delta = 18$ the search restricted to $K(4, 18; q_1^+)$ required 228 hours.

In fact, higher values of δ were treated in a different way by partitioning the search into several tasks that were distributed to several different machines. Using actual timings, we estimated hypothetical timings corresponding to the chip mentioned above. These indicated that the time needed for a complete search for $\delta = 20$ would be about 2700 days, but restricting the search to $K(4, 20; q_1^+)$ reduced this time to about 40 days.

Another feature of our program is its exorbitant redundancy. In an extreme case, a four-dimensional lattice may have 30 points on $\Omega(4, \delta)$, these comprising two point pairs on each of seven facet pairs, and one point pair on the remaining facet pair. When q_1 includes four of these facet pairs, our search over $K(4, \delta, q_1)$ may include the identical lattice sixteen times. Moreover, we might treat each of the 192 lattices in the same equivalence class either eight or sixteen times. All this work might provide a single entry in Table 5. This helps us to understand why the much smaller population space $K(4, \delta, q_1^+)$ often but not always includes at least one of the set of K -optimal lattices associated with the larger (by a factor of up to 24) set. We note that the complexity or the complexity bound would not be affected by this redundancy. It shows itself in the circumstance that an optimal rule was usually found in the first hour of a 100-hour run.

¹Pentium II (Deschutes), 398.13 bogomips processor

For some parts of the search, this redundancy is not important. As an analogy one might compare the task of searching for one of k needles in one haystack with that of searching for one of $100k$ needles in 100 mixed-up haystacks. So long as $k \geq 1$, the time taken to find one needle is to first order the same in either case. If $k = 0$, it takes 100 times as long to complete the search in the second case as in the first case.

APPENDIX: SPECIFICATION OF SETS DEFINED IN THEOREM 3.4

The group \mathcal{G} of coordinate transformations includes transpositions σ_{ij} that interchange coordinates x_i and x_j and reflections ρ_i that replace x_i by $-x_i$. The group can be generated by the four elements σ_{12} , σ_{13} , σ_{14} and ρ_3 , which we have temporarily termed \mathcal{G}_i ($i = 1, 2, 3, 4$). Hence, we can establish the theorem by exploiting the result that $\mathcal{G}_i q \in S_j$ ($i = 1, 2, 3, 4$) whenever $q \in S_j$.

The effect of each of these four transformations on each of the eight facet-pairs is given in the following table.

	Specification	Parity		σ_{12}	σ_{13}	σ_{14}	ρ_3
F_0	+, +, +, +	E	F_0	F_0	F_0	F_0	F_4
F_1	-, +, +, +	O	F_1	F_2	F_4	F_7	F_5
F_2	+, -, +, +	O	F_2	F_1	F_2	F_2	F_6
F_3	-, -, +, +	E	F_3	F_3	F_6	F_5	F_7
F_4	+, +, -, +	O	F_4	F_4	F_1	F_4	F_0
F_5	-, +, -, +	E	F_5	F_6	F_5	F_3	F_1
F_6	+, -, -, +	E	F_6	F_5	F_3	F_6	F_2
F_7	-, -, -, +	O	F_7	F_7	F_7	F_1	F_3

To illustrate the calculation, we confirm the entry for $\sigma_{13}F_3$. By definition, F_3 includes only points of the form $(-a, -b, c, d)$ where a, b, c , and d are individually nonnegative. The corresponding point of $\sigma_{13}F_3$ is obtained by interchange of coordinates 1 and 3, and so is $(c, -b, -a, d)$. Reference to the definition confirms that this point is indeed an element of F_6 . Thirty-two equally trivial calculations will confirm the results presented in this table.

Using these operations, we can transform quartets of facet-pairs into other quartets. We overload the notation above and consider the order of facet-pairs in a quartet as irrelevant; for example, $q(7, 1, 4, 2) = q(1, 2, 4, 7)$.

One way to obtain one of the sets listed below is by constructing a list as follows.

Initially this list contains only one element, in this case perhaps $q_{2a} = q(0, 2, 4, 6)$. At the end of a later stage, it may contain N distinct elements, say, q_1, q_2, \dots, q_N . The next stage comprises calculating $G_i q_k$ for $i = 1, 2, 3, 4$ and $k = 1, 2, \dots, N$, adding these to the list and removing duplicates. If the new list has more than N elements, we proceed to a further stage of the same nature. If the new list has N elements, the same number as in the previous list, we may stop. The current list now comprises a complete list of the elements of S_{2a} .

Again, we illustrate one of these calculations by an example. We evaluate $\sigma_{13}q(0, 2, 4, 6)$. We require from the table the facet-pairs $\sigma_{13}F_j$ for $j = 0, 2, 4, 6$. Reference to the column headed σ_{13} of the table shows these to be F_0, F_2, F_1, F_3 , respectively. These facet-pairs comprise $q(0, 2, 1, 3)$, which is the same as $q(0, 1, 2, 3)$.

Each update of the list involves four such calculations for each of the current N members of the list.

- Elements of S_0
 $q(0, 3, 5, 6), q(1, 2, 4, 7)$
- Elements of S_1
 $q(0, 2, 4, 7), q(0, 1, 4, 7), q(0, 1, 2, 7), q(0, 1, 2, 4),$
 $q(0, 3, 4, 6), q(0, 4, 5, 6), q(0, 1, 3, 5), q(0, 2, 3, 6),$
 $q(0, 3, 4, 5), q(0, 1, 3, 6), q(0, 5, 6, 7), q(0, 3, 5, 7),$
 $q(0, 3, 6, 7), q(0, 1, 5, 6), q(0, 2, 5, 6), q(0, 2, 3, 5),$
 $q(3, 4, 5, 6), q(1, 4, 5, 7), q(2, 4, 6, 7), q(1, 2, 3, 4),$
 $q(1, 2, 4, 6), q(1, 2, 4, 5), q(1, 2, 3, 7), q(2, 4, 5, 7),$
 $q(1, 4, 6, 7), q(1, 2, 5, 7), q(1, 2, 6, 7), q(2, 3, 4, 7),$
 $q(1, 3, 4, 7), q(2, 3, 5, 6), q(1, 3, 5, 6), q(3, 5, 6, 7)$
- Elements of S_{2a}
 $q(0, 2, 4, 6), q(0, 1, 4, 5), q(0, 1, 2, 3), q(0, 3, 4, 7),$
 $q(0, 1, 6, 7), q(0, 2, 5, 7), q(2, 3, 4, 5), q(1, 3, 4, 6),$
 $q(1, 2, 5, 6), q(2, 3, 6, 7), q(1, 3, 5, 7), q(4, 5, 6, 7)$
- Elements of S_{2b}
 $q(0, 2, 4, 5), q(0, 1, 4, 6), q(0, 1, 2, 5), q(0, 2, 3, 4),$
 $q(0, 1, 3, 4), q(0, 1, 2, 6), q(0, 4, 6, 7), q(0, 4, 5, 7),$
 $q(0, 1, 3, 7), q(0, 2, 3, 7), q(0, 2, 6, 7), q(0, 1, 5, 7),$
 $q(2, 4, 5, 6), q(1, 4, 5, 6), q(1, 2, 3, 6), q(1, 2, 3, 5),$
 $q(2, 3, 4, 6), q(1, 3, 4, 5), q(2, 5, 6, 7), q(1, 5, 6, 7),$
 $q(2, 3, 5, 7), q(1, 3, 6, 7), q(3, 4, 6, 7), q(3, 4, 5, 7)$

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