# Tuple lattice sieving

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### Abstract

Lattice sieving is asymptotically the fastest approach for solving the shortest vector problem (SVP) on Euclidean lattices. All known sieving algorithms for solving the SVP require space which (heuristically) grows as  $2^{0.2075n+o(n)}$ , where *n* is the lattice dimension. In high dimensions, the memory requirement becomes a limiting factor for running these algorithms, making them uncompetitive with enumeration algorithms, despite their superior asymptotic time complexity.

We generalize sieving algorithms to solve SVP with less memory. We consider reductions of tuples of vectors rather than pairs of vectors as existing sieve algorithms do. For triples, we estimate that the space requirement scales as  $2^{0.1887n+o(n)}$ . The naive algorithm for this triple sieve runs in time  $2^{0.5661n+o(n)}$ . With appropriate filtering of pairs, we reduce the time complexity to  $2^{0.4812n+o(n)}$  while keeping the same space complexity. We further analyze the effects of using larger tuples for reduction, and conjecture how this provides a continuous trade-off between the memory-intensive sieving and the asymptotically slower enumeration.

## 1. Introduction

The SVP aims to find a shortest non-zero vector in a Euclidean lattice, starting from an arbitrary basis of the lattice. Solving SVP is the cost-dominating component to cryptanalyze lattice-based cryptosystems [14, 19]. The currently best known algorithm with proven correctness and complexity bounds [1] requires memory and time  $2^{n+o(n)}$ , where *n* is the lattice dimension. The best known provable algorithm requiring less space than this is due to Kannan [16]. Its space requirement is polynomial, but its running time is  $n^{n/(2e)+o(n)}$  [12, 13], which is asymptotically slower than sieving. In practice, however, the most competitive implementations rely on variants of Kannan's algorithm that are asymptotically slower and whose correctness can only be guaranteed heuristically [4, 7, 9].

Lattice sieving. The first provable lattice sieving algorithm dates back to the work of Ajtai, Kumar and Sivakumar (AKS) [2]. The AKS algorithm has been progressively refined and simplified in a series of works [11, 22, 24], resulting in the ListSieve algorithms of Micciancio and Voulgaris [20]. Currently, the fastest provable variant of lattice sieving runs in time  $2^{2.465n+o(n)}$  and space  $2^{1.325n+o(n)}$  [23] (see [18] for a quantum acceleration).

In practice, heuristic variants of the lattice sieving algorithms are found to be more efficient. Nguyen and Vidick [22] exhibited a version of AKS that can be heuristically argued correct and which requires a running time of  $(4/3)^{n+o(n)} \approx 2^{0.4150n+o(n)}$  and space of  $(4/3)^{n/2+o(n)} \approx 2^{0.2075n+o(n)}$ . Micciancio and Voulgaris [20] later proposed a heuristic variant of their ListSieve algorithm, namely the GaussSieve algorithm. In practice, the GaussSieve seems to perform well compared with the other variants [20]. It has been investigated further in a series of works

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(see, for example, [8]) and its variants were used to solve some lattice challenges (T. Kleinjung, private communication, 2015). In practice, the GaussSieve algorithm is one of the most promising candidates for lattice sieving algorithms.

Recently, nearest neighbor search techniques have been used to accelerate heuristic sieving algorithms further. The technique was first used in the context of lattice sieving by Laarhoven in [17]. Currently, the best variant is due to Becker, Ducas, Gama and Laarhoven [5], which has a time complexity of  $(3/2)^{n/2+o(n)} \approx 2^{0.2925n+o(n)}$  and space complexity  $(4/3)^{n/2+o(n)}$ .

Reducing the space complexity. Note that the upper bound of the space requirement of all the aforementioned sieve-based algorithms is at least  $(4/3)^{n/2+o(n)}$ . It is conjectured (and experimentally observed [20, 22]) that such space is typically consumed during the execution of these algorithms as well. Let us explain where this bound stems from.

Before we get vectors of norms close to the lattice minimum, sieving algorithms involve lattice vectors that seem uniformly distributed on an *n*-dimensional sphere (or super-thin crust of a ball), and create shorter vectors by subtracting two lattice vectors on the sphere that happen to be sufficiently close. Concretely, this occurs when the angle between the two vectors is less than  $\pi/3$ : their difference results in a shorter vector. One way to heuristically obtain the space complexity bound is to observe that if we have a point  $\boldsymbol{v}$  on a sphere, then it covers a fraction  $\sin^n(\pi/3) = (3/4)^{n/2}$  of this sphere of points at angle at most  $\pi/3$  from this point. If we further assume that each point covers a different part of the sphere, and that overlaps are generally small and almost negligible, then we see that we need about  $(4/3)^{n/2} \approx 2^{0.2075n}$  vectors to cover the sphere. With this many covering points, any extra point expects to see an existing nearby covering point; and their difference leads to a short vector.

Contributions. We propose tuple variants for the ListSieve and GaussSieve algorithms, which we call TupleSieve and TupleMinkowskiSieve, whose memory footprint is smaller than  $2^{0.2075n+o(n)}$ . The main idea is to attempt to create shorter vectors by looking at triples, quadruples, and so on, of vectors rather than pairs of vectors. For triples of vectors, we estimate the space complexity by  $2^{0.1887n+o(n)}$ . For quadruples, the space complexity is about  $2^{0.1724n+o(n)}$ . For growing k (which remains small compared with n), the space complexity seems to scale as  $k^{n/k(1+o(1))}$ , while the running time seems to scale as  $k^{n(1+o(1))}$ . We conjecture that TupleSieve provides a continuous trade-off between the memory-intensive sieving algorithms and the asymptotically slower Kannan enumeration algorithm.

The time complexity of TupleSieve grows very fast with the size k of tuples. If implemented naively, the algorithm has a time complexity which is the kth power of its space complexity. We show that this naive complexity can be reduced by a filtering of pairs of vectors, to remove those that are too unlikely to be extended to a useful triple. More concretely, in the case of k = 3, a triple is useful if either its underlying pairs are useful or if one pair is not too close to orthogonal and the third vector is close to that pair difference. The underlying thought is that if two vectors are almost orthogonal, then a third vector that is far away from both vectors is unlikely to lead to a triple reduction. For k = 3, filtering allows us to decrease the  $2^{0.5661n+o(n)}$ running time of the naive algorithm to  $2^{0.4812n+o(n)}$ .

On removing the heuristics. The correctness of the tuple lattice sieving algorithms presented in this paper is heuristic, as is their complexity analysis. These heuristics are backed by experiments in § 6, but it would be preferable to also better apprehend them in theory.

The main obstacle towards proving correctness of sieving algorithms is that the vector combinations may all result in the  $\mathbf{0}$  vector after some stage, leading us to miss shortest non-zero vectors. This difficulty is typically circumvented by adding perturbations to the vectors to 'hide' the lattice structure to the algorithm (see, for example, [20]). We believe that such a technique could also be applied to our algorithms, although with a significant cost increase.

The problem of obtaining rigorous bounds on the space complexity is, in our opinion, much more challenging (and mathematically enticing). Bounding the space complexity of ListSieve and GaussSieve can be reformulated in terms of spherical codes: how large can a list of points on the unit sphere be if we assume that all pairs of points have angle at least  $\pi/3$  (that is, their difference has norm >1)? In the case of k-tuple sieving, the list of points on the unit sphere is such that any sum of at most k list vectors has norm >1. As the constraint gets stronger with increasing k, the maximum list size cannot increase. Can it be shown that it decreases?

Road-map. Section 2 gives preliminaries on the geometry of lattices. Section 3 presents the TupleSieve and its complexity analysis. Section 4 describes the filtered TupleSieve, which is designed to optimize its running time. The TupleSieve is a generalization of ListSieve. A GaussSieve-like variant of the TupleSieve, called TupleMinkowskiSieve, is discussed in § 5: similarly to GaussSieve and ListSieve, TupleMinkowskiSieve is more complex to study than the TupleSieve but, in practice, has more potential. Finally, experimental results are discussed in § 6: we test heuristics used in the theoretical analysis of the TupleSieve and we test the practical efficiency of TupleMinkowskiSieve.

#### 2. Preliminaries

First, we introduce some notation and recall some elementary geometric facts that we will use when analyzing tuple lattice sieving algorithms.

Let  $\mathcal{B}(\boldsymbol{v},r)$  be the ball of radius r around  $\boldsymbol{v} \in \mathbb{R}^n$ . Denote, in short,  $\mathcal{B}(\boldsymbol{v}) = \mathcal{B}(\boldsymbol{v},1)$  and  $\mathcal{B} = \mathcal{B}(\mathbf{0},1)$ . We let  $\mathcal{S}$  denote the unit sphere. Let us write  $|\Omega|$  for the volume of a (measurable) set  $\Omega \subset \mathbb{R}^n$ . For a set  $\Omega$  of finite measure, we let  $U(\Omega)$  denote the uniform distribution on  $\Omega$ . For two functions A, B of n, we write  $A \propto B$  if there exist two constants c and c' such that  $A \leq n^c \cdot B$  and  $B \leq n^{c'} \cdot A$  for large n.

# 2.1. Geometric properties of the unit sphere

We first recall the following simple geometric observation, regarding the norm of the difference between two vectors on the sphere.

LEMMA 2.1 (Cosine law). For vectors  $v_1, v_2$  with angle  $\theta$ ,

$$\|v_1 - v_2\|^2 = \|v_1\|^2 + \|v_2\|^2 - 2\|v_1\|\|v_2\|\cos\theta.$$

Thus for two unit vectors  $v_1, v_2 \in S$  with angle  $\theta, ||v_1 - v_2|| = \sqrt{2(1 - \cos \theta)}$ .

Throughout the paper, we are interested in the covering of the unit sphere S by unit balls  $\mathcal{B}(\boldsymbol{v})$ : how many vectors  $\boldsymbol{v}$  are needed such that the union of several balls  $\mathcal{B}(\boldsymbol{v})$  covers (most of) S? The following lemma considers the probability mass of different portions of the unit sphere, when considering the uniform distribution over the sphere.

LEMMA 2.2. The density function  $f(\theta)$  of the angle  $\theta \in [0, \pi/2)$  between any fixed vector in S and a vector sampled independently from U(S) satisfies  $f(\theta) \propto (\sin \theta)^n$ .

A spherical cap of height h in a ball of radius r is any set that may be obtained by applying an isometry to  $\{x \in \mathcal{B}(\mathbf{0}, r) : x_n \ge r - h\}$ . We let C(h, r) denote the volume of a spherical cap with parameters h and r.

LEMMA 2.3 [20, Lemma 4.1]. A spherical cap of height h and radius r has volume

$$C(h,r) \propto (r^2 - (r-h)^2)^{n/2} \cdot |\mathcal{B}|.$$

The following lemma will be useful for estimating the part of the sphere that is covered by a (unit) ball at arbitrary distance from the origin.

LEMMA 2.4. Two balls  $\mathcal{B}(\boldsymbol{v}_1, r_1)$  and  $\mathcal{B}(\boldsymbol{v}_2, r_2)$  at distance  $\|\boldsymbol{v}_1 - \boldsymbol{v}_2\| = d$  and radii  $r_1, r_2$  such that  $r_1, r_2 < d < r_1 + r_2$  satisfy

$$|\mathcal{B}(\boldsymbol{v}_1, r_1) \cap \mathcal{B}(\boldsymbol{v}_2, r_2)| \propto \left(rac{-d^4 + 2d^2(r_1^2 + r_2^2) - (r_1^2 - r_2^2)^2}{4d^2}
ight)^{n/2} \cdot |\mathcal{B}|.$$

In particular,  $|\mathcal{B}(\boldsymbol{v}_1) \cap \mathcal{B}(\boldsymbol{v}_2)| \propto (1 - d^2/4)^{n/2} \cdot |\mathcal{B}|.$ 

Proof. Without loss of generality, we assume that  $\mathbf{v}_1 = \mathbf{0}$  and  $\mathbf{v}_2 = d\mathbf{e}_1$ . The intersection of these two balls is partitioned in two spherical caps. Let  $\mathbf{a}$  be a point at distance  $r_1$  from  $\mathbf{0}$  and distance  $r_2$  from  $d\mathbf{e}_1$  and consider the triangle formed by  $\mathbf{0}, \mathbf{a}$  and  $d\mathbf{e}_1$ . This triangle has sides  $d, r_1, r_2$ . To find  $|\langle \mathbf{a}, d\mathbf{e}_1 \rangle|$ , we use the law of cosines in the point  $\mathbf{0}$  to establish that this angle in the triangle satisfies  $\cos \phi = (r_1^2 + d^2 - r_2^2)/(2r_1d)$ . Using the cosine definition, we then know that  $\cos \phi = |\langle \mathbf{a}, d\mathbf{e}_1 \rangle|/r_1$ . We define  $x = |\langle \mathbf{a}, d\mathbf{e}_1 \rangle| = (r_1^2 + d^2 - r_2^2)/(2d)$ . The spherical cap associated to the ball of radius  $r_1$  has volume

$$C(r_1 - x, r_1) \propto \left(\frac{-d^4 + 2d^2(r_1^2 + r_2^2) - (r_1^2 - r_2^2)^2}{4d^2}\right)^{n/2} \cdot |\mathcal{B}|.$$

The other spherical cap has the same volume  $C(r_2 - (d - x), r_2) \propto C(r_1 - x, r_1)$ . As the total volume of the intersection is the sum of these two values, the result follows.

### 2.2. Euclidean lattices

A (full-rank) lattice of  $\mathbb{R}^n$  is the set  $\mathcal{L}[(\mathbf{b}_i)_i] = \sum_i \mathbb{Z} \mathbf{b}_i \subset \mathbb{R}^n$  of all integer combinations of some *n* linearly independent vectors  $(\mathbf{b}_i)_{i \leq n}$  of  $\mathbb{R}^n$ . In this set-up, the vectors  $(\mathbf{b}_i)_{i \leq n}$  are said to form a basis of  $\mathcal{L}[(\mathbf{b}_i)_i]$ . Note that any given lattice of dimension  $n \geq 2$  admits an infinite number of bases. A lattice  $\mathcal{L}$  contains shortest non-zero vectors; their common norm is referred to as the lattice minimum and denoted by  $\lambda(\mathcal{L})$ . The algorithmic task of finding a shortest non-zero vector of  $\mathcal{L}[(\mathbf{b}_i)_i]$  given  $(\mathbf{b}_i)_i$  as input is known as the shortest vector problem (SVP).

In tuple variants of the GaussSieve algorithm (TupleMinkowskiSieve), we consider a list of lattice vectors which are tuple-wise Minkowski reduced.

DEFINITION 1 (Minkowski reduction). A basis  $(\mathbf{b}_i)_{i \leq n}$  of a lattice  $\mathcal{L}$  is Minkowski reduced if the  $\mathbf{b}_i$  are sorted by non-decreasing norms and if, for all i < n, the vector  $\mathbf{b}_i$  has minimal norm among all vectors  $\mathbf{b} \in \mathcal{L}$  such that  $(\mathbf{b}_1, \mathbf{b}_2, \dots, \mathbf{b}_{i-1}, \mathbf{b})$  can be extended to a basis of  $\mathcal{L}$ .

Note that Gauss reduction is Minkowski reduction in the special case of n = 2.

# 3. Tuple sieving

Let us first describe the general strategy of sieving algorithms. These algorithms start with a large number of long lattice vectors, for example, created by sampling from a discrete Gaussian over the lattice with a large standard deviation [10]. They try to gradually reduce their norms by considering simple combinations. The process of combining vectors to create shorter lattice vectors is the sieving procedure, which produces shorter and shorter lattice vectors when applied iteratively; vector combinations whose norm is above some threshold are discarded during the sieving procedure, while short combinations are kept for the next iteration.

To make sure that we are not losing too many vectors during this sieving procedure and to bound the running time, we use a 'covering' set which provably cannot become too large (for example, the center set in the Nguyen and Vidick sieve [22], or the list in the Micciancio and Voulgaris sieve [20]). In general, the vectors in this 'covering' set are not too close to each other. If two current vectors are sufficiently close, we will reduce them before putting them into this 'covering' set.

To prove correctness (that is, that shortest non-zero vectors are not avoided in the successive sieves), the standard approach is using perturbations: one adds a small error vector to each lattice vector, so that the lattice is blurred from the perspective of the sieves. This allows us to argue that the probability of finding a non-zero vector cannot be arbitrarily small compared with the probability of getting the zero vector. We refer to [20] for more details.

To ease the analysis, we consider a simplified algorithm named DoubleSieve (see Algorithm 1) and we do not consider these perturbations, which appear to be an artefact of the proof of correctness, rather than a procedure necessary to make sure that the algorithm succeeds.

Algorithm 1 DoubleSieve	Algorithm 2 TripleSieve
1: $L' \leftarrow \{\}$	1: $L' \leftarrow \{\}$
2: for each $\boldsymbol{v}, \boldsymbol{w} \in \mathrm{L}$ do	2: for each $oldsymbol{v},oldsymbol{w},oldsymbol{x}\in \mathrm{L}$ do
3: <b>if</b> $\  \boldsymbol{v} \pm \boldsymbol{w} \  \leqslant \gamma \cdot R$ then	3: <b>if</b> $\  \boldsymbol{v} \pm \boldsymbol{w} \pm \boldsymbol{x} \  \leqslant \gamma \cdot R$ then
4: $\mathrm{L}' \leftarrow \mathrm{L}' \cup \{ \boldsymbol{v} \pm \boldsymbol{w} \}^{\dagger}$	4: $\mathbf{L}' \leftarrow \mathbf{L}' \cup \{ \boldsymbol{v} \pm \boldsymbol{w} \pm \boldsymbol{x} \}^{\dagger}$

The DoubleSieve follows the same main procedure as the other sieves. In short, the main point is to reduce pairs of vectors in L to get a new list L'. The norms of the vectors in L are bounded by some constant R, and the norms of the vectors in L' are then bounded by  $\gamma \cdot R$  for some geometric factor  $\gamma < 1$ . This factor  $\gamma$  ensures that, in each iteration of the sieve, we make some progress with reducing the norms of the list vectors. In the analysis, we will assume that  $\gamma \approx 1$ , so that we do not need many more points to create a large output list. Note that we can choose  $\gamma = 1 - 1/\text{poly}(n)$ , while ensuring that the total number of sieves remains poly(n): indeed, we may start with an initial radius  $R \leq 2^n \cdot \lambda(\mathcal{L})$ , by pre-processing the input basis with the LLL lattice reduction algorithm (Lenstra–Lenstra–Lovász), and stop with a final radius  $\gamma^{\text{poly}(n)}R \approx \lambda(\mathcal{L})$ .

Note that a sieving step aims at reducing the norms of the current vectors by a multiplicative factor  $\gamma > 1$ , in order to prevent the whole procedure from stalling. As a result, perturbation-free sieving algorithms (and our extensions) are unlikely to solve the SVP and seem limited to finding non-zero lattice vectors of norms  $\leq t \cdot \lambda(\mathcal{L})$  for some t that is close to one. We make do with only finding near shortest non-zero lattice vectors.

An extension (TripleSieve) of this DoubleSieve algorithm is presented in Algorithm 2 above, where, instead of pairs of vectors, we consider triples to find shorter lattice vectors. Note that for pairs of vectors, considering combinations  $v \pm w$  is the best one can do: if there exists some integer linear combination  $z_1v + z_2w$  which has norm < R for vectors v, w of norms R, then also one of the vectors  $v \pm w$  must have norm < R. For triples (or tuples) it is harder to tell which finite set of linear combinations must be considered to guarantee obtaining the strongest notion of reduction in our list. The Minkowski conditions described in [21, 26] show that, for triples of vectors, it suffices to consider combinations  $v \pm w, v \pm x, w \pm x$  and  $v \pm w \pm x$ for reduction to achieve Minkowski reducedness for a triple of lattice vectors  $\{v, w, x\}$ . If we assume that **0** belongs to L, then these are exactly the combinations considered in Algorithm 2.

#### 3.1. Cost analysis of the DoubleSieve

We now proceed with a sketch of the cost analysis for the Double Sieve, which will serve as a guideline for analyzing the TripleSieve in  $\S 3.2$ .

 $<sup>^{\</sup>dagger} \mathrm{The}$  sign choice is the same as in the step above.

Suppose that we start with a certain list of size  $|\mathbf{L}| = N$  and let  $\boldsymbol{v}, \boldsymbol{w} \in \mathbf{L}$ . Let both vectors have norm approximately R; significantly shorter vectors are also shorter than  $\gamma R$  and are immediately added to L'. Now the condition  $\|\boldsymbol{v} - \boldsymbol{w}\| < \gamma R$  for  $\gamma \approx 1$ , equivalently, corresponds to  $\boldsymbol{w} \in \mathcal{B}(\mathbf{0}, R) \cap \mathcal{B}(\boldsymbol{v}, \gamma R)$ . To estimate the probability of finding a vector  $\boldsymbol{v} - \boldsymbol{w}$  satisfying  $\|\boldsymbol{v} - \boldsymbol{w}\| < \gamma R$ , we use the following heuristic assumption, introduced in [22].

HEURISTIC 1. We assume that each time DoubleSieve is called, the vectors v/||v|| for  $v \in L$  are independent and identically distributed uniformly distributed points on the unit sphere.

We may restrict ourselves to analyzing the DoubleSieve for listing vectors of near-identical norms (as they are coming from a prior sieve, and if they were much shorter than expected, they could have been kept for later sieves). Further, as the DoubleSieve is scale invariant, we may restrict the study to the case where L consists of unit vectors.

Assuming Heuristic 1 holds, we can now estimate the probability that  $\boldsymbol{w} \in \mathcal{B} \cap \mathcal{B}(\boldsymbol{v},\gamma)$ by the relative mass of the corresponding spherical cap on the sphere. Letting  $\gamma \approx 1$  and  $\|\boldsymbol{v}\| \approx 1$ , this probability is  $p \approx \sin^n(\pi/3) = (3/4)^{n/2}$ . So, given any lattice vector in the list, the probability that a second vector is going to lead to a good combination which can be used for L' is proportional to  $(3/4)^{n/2}$ . Alternatively, one could say that each vector in the list covers a fraction  $(3/4)^{n/2}$  of the sphere; vectors falling inside this spherical cap will lead to pairwise reductions, while vectors outside will not. Assuming that the intersections of these spherical caps are negligible, we therefore approximately need  $1/p \approx (4/3)^{n/2}$  points to cover the entire sphere: using a list L of size poly $(n) \cdot (4/3)^{n/2}$  guarantees that with overwhelming probability; any other vector can be reduced with one of the list vectors, while, if L is significantly smaller than  $(4/3)^{n/2}$ , then, with overwhelming probability, a random lattice vector is not covered by this list L, meaning that we will lose many points in each iteration of the sieve.

To summarize, the crucial equation for the list size N to guarantee that 1-o(1) of the sphere is covered with this list is given by

$$N \cdot \left(\frac{3}{4}\right)^{n/2} \ge 1 - o(1).$$

Finally, by taking, for instance,  $\gamma = 1 - 1/n$ , it is guaranteed that only a polynomial number of iterations is needed to go from an initial list of long lattice vectors to a list of vectors of norm at most  $\lambda(\mathcal{L})$ . The time complexity is therefore dominated by poly(n) applications of the DoubleSieve, whose cost is quadratic in the list size N. This leads to a memory complexity of  $N \propto (4/3)^{n/2}$  and a time complexity of  $N^2 \propto (4/3)^n$ .

### 3.2. Cost analysis of the TripleSieve

In the TripleSieve, not only single list vectors cover subsets of the sphere, but also sums and differences of pairs of list vectors cover parts of the sphere; if  $\boldsymbol{v} - \boldsymbol{w} - \boldsymbol{x}$  is short, then  $\boldsymbol{x}$  is covered by the spherical cap  $\mathcal{B}(\boldsymbol{0},R) \cap \mathcal{B}(\boldsymbol{v} - \boldsymbol{w},\gamma R)$ . So not only do the N single vectors cover part of the sphere, but also the roughly  $N^2$  sums and differences of list vectors  $\boldsymbol{v} \pm \boldsymbol{w}$  each cover a region of the sphere in the sense that any vector  $\boldsymbol{x}$  in one of these regions can be reduced with  $(\boldsymbol{v} \pm \boldsymbol{w})$ . Intuitively, this explains why fewer vectors will be needed to cover the entire sphere (and to guarantee that L' will not be shorter than L).

In the analysis below, we will make use of the following generalization of Heuristic 1.

HEURISTIC 2. We assume that each time the TripleSieve is called, the vectors  $\boldsymbol{v}/\|\boldsymbol{v}\|$  for  $\boldsymbol{v} \in L$  are independent and identically distributed uniformly distributed points on the unit sphere and that all vectors  $\boldsymbol{v} \in L$  and  $\boldsymbol{v} \pm \boldsymbol{w}$  for  $\boldsymbol{v} \neq \boldsymbol{w} \in L$ , with sign choice minimizing the norm, behave as if they were independent.

Note that the vectors  $v \pm w$  for  $v \neq w \in L$  cannot be independent as they are deterministically obtained from a much smaller set of points. A heuristic stating that they are independent would hence be invalid. However, we still we may assume that this non-independence does not have an impact on the analysis below.

As in the analysis of DoubleSieve, we assume that the vectors of L all lie on the unit sphere S. Now the vectors  $v \pm w$  generally do not lie on S and the part of S that is covered by  $\pm v \pm w$  depends exactly on the norm of  $v \pm w$ . Let us express this norm in terms of the angle  $\theta \in [0, \pi/2]$  between v and w. For unit vectors v and w,

$$\min \|\boldsymbol{v} \pm \boldsymbol{w}\|^2 = 2(1 - \cos \theta).$$

Next, note that if a vector has norm r, then the part of the unit sphere it covers is proportional to  $(1 - r^2/4)^{n/2}$  (by Lemma 2.4). Therefore, if  $\boldsymbol{v}$  and  $\boldsymbol{w}$  have angle  $\theta$ , then the vector  $\boldsymbol{v} \pm \boldsymbol{w}$  covers a fraction of the spherical surface equal to at least

$$g(\theta) = (1 - \| \boldsymbol{v} \pm \boldsymbol{w} \|^2 / 4)^{n/2} = \cos(\theta / 2)^n$$

Here we used the half-angle identity  $\cos^2(\phi/2) = (1 + \cos \phi)/2$  that holds for arbitrary  $\phi$ .

We now compute the expected portion of S that is covered by the difference vector between two list vectors. By Lemma 2.2, the density of angles between pairs of vectors is proportional to  $f(\theta) \propto \sin(\theta)^n$ . The expected value of the part of S covered by a pair of vectors is therefore

$$\mathbb{E}_{\theta}[g(\theta)] \propto \int_{0}^{\pi/2} f(\theta)g(\theta) \, d\theta = \int_{0}^{\pi/2} \left(\sin(\theta)\cos\left(\frac{\theta}{2}\right)\right)^{n} d\theta.$$

Note that the integrand is exponential in n, and so the asymptotic scaling of the entire integral is determined by the maximum value of the integrand. Ignoring polynomial terms,

$$\mathbb{E}_{\theta}[g(\theta)] \propto \left( \max_{\theta} \sin(\theta) \cos\left(\frac{\theta}{2}\right) \right)^n.$$

With some elementary trigonometric manipulation we see that this maximum is attained at  $\theta = \arccos(1/3)$  and, in this point, the function takes value  $4/(3\sqrt{3}) = \sqrt{16/27}$ . So we obtain

$$\mathbb{E}_{\theta}[g(\theta)] = \left(\frac{16}{27}\right)^{n/2}.$$

To figure out how large the list size must be to cover the entire sphere, suppose we have N points in our list. Then, to guarantee that the single points and pairs of points in the list together cover a fraction 1 - o(1) of the sphere, assuming that the overlap between these regions is asymptotically negligible, this leads to the condition on N

$$N \cdot \left(\frac{3}{4}\right)^{n/2} + N^2 \cdot \left(\frac{16}{27}\right)^{n/2} \ge 1 - o(1).$$

The first term corresponds to the area covered by single list vectors, whereas the second term stems from pairs of list vectors. If N is much smaller than  $(4/3)^{n/2}$ , then the first term is exponentially smaller than 1 and the second term dominates<sup>†</sup>. So solving for N in  $N^2 \cdot (16/27)^{n/2} \propto 1$ , we obtain  $N \propto (27/16)^{n/4} \approx 2^{0.1887n}$ . Note that this list size N is strictly smaller than the DoubleSieve list size of  $\propto (4/3)^{n/2} \approx 2^{0.2075n}$ .

Finally, similar to that of the DoubleSieve, the cost of this algorithm is dominated by having to store the lists of vectors (memory) and having to consider all pairs/triples of vectors for the sieve (time). This directly leads to heuristic space and time complexities of  $(27/16)^{n/4} \approx 2^{0.1887n}$  and  $(27/16)^{3n/4} \approx 2^{0.5662n}$  for a naive cubic search over all triples.

<sup>&</sup>lt;sup>†</sup>The imbalance of the two terms implies that in the TripleSieve we may consider only combinations of triples of vectors and forget about combinations of pairs of vectors.

#### 3.3. Cost analysis of the k-TupleSieve

Algorithm 3 generalizes the DoubleSieve and the TripleSieve to arbitrary k-tuples. The DoubleSieve and the TripleSieve correspond to setting k = 2 and k = 3, respectively. Similarly to the DoubleSieve and the TripleSieve, the sign choice at Step 4 is identical to that of Step 3. Further, we may assume that vector **0** belongs to L so that the test of Step 3 actually allows to consider combinations of up to k vectors. Or we may argue (thanks to the analysis below) that only combinations of exactly k (non-zero) list vectors are likely to lead to norm reductions, and discard combinations of fewer vectors.

Al	gorithm 3 k-TupleSieve
1:	$L' \leftarrow \{\}$
2:	for each $oldsymbol{v}_1,oldsymbol{v}_2,\ldots,oldsymbol{v}_k\in\mathrm{L}$ do
3:	$\mathbf{if}  \left\  \boldsymbol{v}_1 \pm \boldsymbol{v}_2 \pm \ldots \pm \boldsymbol{v}_k \right\  \leqslant \gamma \cdot R  \mathbf{then}$
4:	$\mathrm{L}' \leftarrow \mathrm{L}' \cup \{ oldsymbol{v}_1 \pm oldsymbol{v}_2 \pm \ldots \pm oldsymbol{v}_k \}$

We now generalize the previous analyses to tuple sieving. Let k be fixed and let  $v_1, \ldots, v_{k-1}$  be independently sampled from U(S). Define  $v'_1 = v_1$  and, for  $1 < i \leq k$ , let  $v'_i$  be either  $v_i$  or  $-v_i$ , so that  $\langle v'_i, \sum_{j < i} v'_j \rangle \leq 0$ . Note that the angles  $\theta_i$  between  $\sum_{j < i} v'_j$  and  $v_i$  are independent and have densities proportional to  $(\sin \theta)^n$ , by Lemma 2.2. Further, observe that  $\theta_2, \ldots, \theta_i$  fully determine  $f_i := \|\sum_{j \leq i} v'_j\|$ . Indeed,  $f_1 = 1$  and, for i > 1,

$$f_i^2 = f_{i-1}^2 - 2f_{i-1}\cos\theta_i + 1.$$

Now, the proportion  $g_{k-1}$  of S at distance  $\leq 1$  from  $\sum_{i < k} v'_i$  covered by this sum is proportional to the size of the intersection of two unit balls centered at **0** and at a point at distance  $f_{k-1}$  from **0**. By Lemma 2.4,  $g_{k-1} \propto (1 - f_{k-1}^2/4)^{n/2}$ . Hence

$$\alpha_{k-1} := \mathbb{E}_{\boldsymbol{v}_1, \dots, \boldsymbol{v}_{k-1}}[g_{k-1}] \propto \left(\max_{\theta_2, \dots, \theta_{k-1}} \sqrt{1 - \frac{f_{k-1}^2}{4}} \prod_{1 < i < k} \sin \theta_i\right)^n.$$
(3.1)

By the above, a list size L satisfying

$$\alpha_1^n L + \alpha_2^n L^2 + \ldots + \alpha_{k-1}^n L^{k-1} = 1$$

suffices to cover the sphere by k-tuples. The space requirement is  $S_k = (\max_{i < k} \alpha_i^{1/i})^n$  and the time requirement  $T_k$  is  $S_k^k$ . Note that  $\alpha_1 = \sqrt{4/3}$  and  $\alpha_2 = \sqrt{27/16}$ , as in the previous subsections.

We give estimates for the triple and quadruple sieves. For k = 3, we have previously seen that the target function was

$$\left(\sqrt{1-\frac{f_2^2}{4}}\right)\sin\theta_2 = \left(\sqrt{\frac{1}{2}\cos(\theta_2) + \frac{1}{2}}\right)\sin(\theta_2) = \cos\left(\frac{\theta_2}{2}\right)\sin(\theta_2).$$

The maximum is achieved at  $\theta_2 = \arccos(1/3)$  and the maximum value is  $4/(3\sqrt{3})$ , leading to a list size  $2^{0.1887n+o(n)}$  for the TripleSieve. For k = 4, the target function is

$$\sqrt{\frac{1}{2}\sqrt{2-2\cos(\theta_2)}\cos(\theta_3) + \frac{1}{2}\cos(\theta_2) + \frac{1}{4}\sin(\theta_2)\sin(\theta_3)}.$$

Numerically optimizing over  $\theta_2$  and  $\theta_3$ , we obtain a quadruple sieve list size  $2^{0.1724n+o(n)}$ .

For even larger tuple sizes k, Table 1 lists the numerical values for the space complexity of the k-TupleSieve for tuple sizes k from 3 to 15. The first column denotes the k-tuples used. The second column  $(\log_2|\mathbf{L}|)/n$  gives the list size estimate for the k-tuples. The columns ' $\cos \theta_i$ ' denote the cosines for the optimized angles for the target function in equation (3.1). We used

function MINIMIZE in SAGEMATH [27] to optimize equation (3.1), and we obtained equivalent results using FINDMINIMUM in MATHEMATICA [28].

TABLE 1. Estimates of list size for k-tuple sieving.

k	$\frac{\log_2( \mathbf{L} )}{n}$	$\cos \theta_2$	$\cos \theta_3$	$\cos \theta_4$	$\cos \theta_5$	$\cos \theta_6$	$\cos \theta_7$	$\cos \theta_8$	$\cos  heta_9$	$\cos \theta_{10}$	$\cos \theta_{11}$	$\cos \theta_{12}$	$\cos \theta_{13}$	$\cos \theta_{14}$
3	0.1887	0.333												
4	0.1724	0.250	0.408											
5	0.1587	0.200	0.316	0.447										
6	0.1473	0.167	0.258	0.354	0.471									
7	0.1376	0.143	0.218	0.293	0.378	0.488								
8	0.1293	0.125	0.189	0.250	0.316	0.395	0.500							
9	0.1221	0.111	0.167	0.218	0.272	0.333	0.408	0.509						
10	0.1158	0.100	0.149	0.194	0.239	0.289	0.346	0.418	0.516					
11	0.1102	0.091	0.135	0.174	0.213	0.255	0.302	0.357	0.426	0.522				
12	0.1052	0.083	0.123	0.158	0.192	0.228	0.267	0.312	0.365	0.433	0.527			
13	0.1007	0.077	0.113	0.145	0.175	0.207	0.240	0.277	0.320	0.372	0.439	0.531		
14	0.0967	0.071	0.105	0.134	0.161	0.189	0.218	0.250	0.286	0.327	0.378	0.443	0.534	
15	0.0930	0.067	0.098	0.124	0.149	0.174	0.200	0.228	0.258	0.293	0.333	0.383	0.447	0.538

#### 3.4. Conjectured large-k asymptotics

Finally, we conclude this theoretical analysis with conjectured large-k asymptotics of the time and space complexities of the k-TupleSieve. For small k = 2, 3 we have exact algebraic expressions for the heuristic space complexities and, if we attempt to match similar expressions to the numeric data in the first column of Table 1 using the Inverse Symbolic Calculator [15], we obtain a conjectured general expression for the heuristic space complexity for arbitrary k.

CONJECTURE. The heuristic asymptotic space complexity |L| of the k-TupleSieve satisfies

$$L|^{1/n} = \frac{k^{k/(2k-2)}}{\sqrt{k+1}}.$$

This formula matches our numerically obtained results for k = 2, ..., 15 up to the first 50 digits. We plot the numerical results and the conjectured space complexity curve in Figure 1. Note that, indeed, for k = 2, 3 this formula gives  $|L|^{1/n} = \sqrt{4/3}$  and  $|L|^{1/n} = (27/16)^{1/4}$ .

Assuming that this formula is correct, we can study the limiting behavior of large tuples. First, observe that the expression above scales as  $k^{1/k+o(1/k)}$ . In other words, the list size asymptotically scales as  $|\mathbf{L}| = k^{n/k+o(k)}$  and the corresponding time complexity is given by  $k^{n+o(n)}$ . If we let k approach n, then we see that the estimated list size approaches  $|\mathbf{L}| \to n^{1+o(1)}$ , while the time complexity scales as  $n^{n+o(n)}$ . This conjectured asymptotic scaling matches (up to constants in the exponents) the complexities of Kannan's enumeration algorithm [16]. Tuple lattice sieving could therefore be considered a way to obtain a continuous trade-off between the asymptotically fast but memory-intensive heuristic sieving algorithms (small k), and the memory-efficient and asymptotically slow enumeration methods (large k).

#### 4. Filtered triple sieving

In the analysis of the TripleSieve above, we saw that vectors with angle  $\theta$  significantly smaller than  $\arccos(1/3)$  lead to a lot of the sphere being covered, but these vectors do not appear often (thus in the algorithm, we should try to use them all); vectors at angle  $\theta \approx \arccos(1/3)$ 

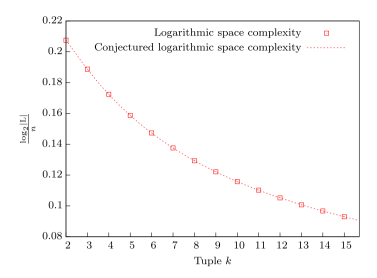


FIGURE 1. The numerically obtained heuristic logarithmic space complexities for k-tuple sieving up to k = 15 and the conjectured pattern for arbitrary k from (3.1).

cover a decent amount of the sphere and contribute 1 - o(1) of the found collisions among triples of vectors; and vectors at angle  $\theta$  significantly larger than  $\arccos(1/3)$  appear very often, but rarely lead to reductions, even though this case appears very often (thus, in the algorithm, we can ignore them). This motivates the filtered TripleSieve described in Algorithm 4.

Alg	Algorithm 4 Filtered TripleSieve							
1:	$L' \leftarrow \{\}$							
2:	$ ext{for each } oldsymbol{v}, oldsymbol{w} \in \operatorname{L} \operatorname{\mathbf{do}}$							
3:	$\mathbf{if} \;  \langle oldsymbol{v}, oldsymbol{w}  angle  \geqslant rac{1}{3} \; \mathbf{then}$							
4:	$\mathbf{for} \; \mathbf{each} \; \overset{{}_\circ}{\boldsymbol{x}} \in \mathrm{L} \; \mathbf{do}$							
5:	$\mathbf{if}  \left\  \boldsymbol{v} \pm \boldsymbol{w} \pm \boldsymbol{x} \right\  \leqslant \gamma R  \mathbf{then}$							
6:	$\mathrm{L}' \gets \mathrm{L}' \cup \{ \bm{v} \pm \bm{w} \pm \bm{x} \}$							

In this extension of the TripleSieve, the third search over the list is only performed if the first two vectors are sufficiently close. Asymptotically, this means that the same number of good triples are still found, but this significantly reduces the cost of the algorithm: after all, out of all pairs of vectors, only those with pairwise angle less than  $\theta_1 = \arccos(1/3)$  survive. The fraction of pairs that survive is proportional to  $(\sin \theta_1)^n$  or, after a trigonometric exercise, a fraction  $p = (2\sqrt{2}/3)^n$  of all pairs of vectors survive the first round. The time cost of the algorithm is then  $N^2(1 + p \cdot N)$ , which, as N is much larger than 1/p, leads to the following result.

PROPOSITION 4.1. Under the aforementioned heuristic assumptions, the filtered triple sieve solves SVP in time  $(27/16)^{3n/4} \cdot (2\sqrt{2}/3)^n = 2^{0.4812n}$  and space  $(27/16)^{n/4} = 2^{0.1887n}$ .

Note that such a filtering strategy may also be applied to the k-TupleSieve with larger k.

# 5. Tuple MinkowskiSieve

In the previous sections, we described the tuple variants of the ListSieve algorithm. In practice, it has been observed that the GaussSieve algorithm of Micciancio and Voulgaris [20] is more

efficient (in terms of both time and space) than the ListSieve algorithm. Hence, for practical considerations, we devise a tuple-like GaussSieve algorithm.

# 5.1. GaussSieve

We recall the GaussSieve algorithm of Micciancio and Voulgaris [20]. The philosophy of the algorithm is to make every pair of distinct vectors in a list Gauss reduced.

In GaussSieve, we first set up and maintain a list L of vectors and a stack S of vectors in the algorithm. They are initially empty. To begin the GaussSieve algorithm, we sample a new vector  $\boldsymbol{p}$ . We add  $\boldsymbol{p}$  to the list L. The list L consists of lattice vectors that are always pairwise Gauss reduced and this property is maintained during the execution of the algorithm. For each new sampled vector  $\boldsymbol{p}$ , we may modify  $\boldsymbol{p}$  (to  $\boldsymbol{p}'$ ) and the existing vectors in the list L (to L') so that  $L' \cup \boldsymbol{p}'$  is pairwise Gauss reduced. Therefore, for each  $\boldsymbol{p}$ , we not only reduce  $\boldsymbol{p}$  (to  $\boldsymbol{p}'$ ) using the vectors  $\boldsymbol{v} \in L$ , but also reduce the vectors  $\boldsymbol{v} \in L$  using  $\boldsymbol{p}'$ . If  $\boldsymbol{p}$  (or  $\boldsymbol{p}'$ ) is equal to some  $\boldsymbol{v} \in L$ , then we discard it and count it as a collision of zero vectors. If some list vector  $\boldsymbol{v} \in L$  has been reduced by  $\boldsymbol{p}$ , then the list itself may not be pairwise Gauss reduced any more (since  $\boldsymbol{v}$  might be modified). We then move the modified vector  $\boldsymbol{v}'$  to the stack S and attempt to reduce it in the following reductions (taking  $\boldsymbol{v}'$  as a new sample  $\boldsymbol{p}$ ).

Note that, in GaussSieve, we are not Gauss reducing the two vectors in a one-off way. Instead, we only reduce one vector at one iteration and then put the modified vector in the stack (if it is a vector in the list), and perhaps change it in the future.

In practice, we can terminate the algorithm if we have found a short enough vector or if the number of collisions of zero vectors reach some bound.

## 5.2. Tuple MinkowskiSieve

The Gauss-reduced condition for two vectors can be generalized to the Minkowski-reduced condition for tuple vectors (see Definition 1). The greedy reduction algorithm of [21, 25] can be used to efficiently compute Minkowski-reduced bases for lattices of dimension  $\leq 4$ .

We describe the triple MinkowskiSieve algorithm in Algorithms 5 and 6. The quadruple MinkowskiSieve algorithm can be designed similarly.

Algorithm 5 TRIPLEMINKOWSKISIEVE $((b_i)_i)$	<b>Algorithm 6</b> TRIPLEREDUCE $(p, L, S)$
1: $L \leftarrow \{0\}, S \leftarrow \{\}$ 2: while cond do 3: if S is not empty then 4: $p \leftarrow S.POP()$ 5: else 6: $p \leftarrow SAMPLEGAUSSIAN((b_i)_i)$ 7: $p \leftarrow TRIPLEREDUCE(p, L, S)$ 8: if $p \neq 0$ then insert $p$ to L	1: Loop <sup>†</sup> { $\forall v \in L$ , reduce $p$ by $v$ } 2: if $p = 0$ then Return $p$ 3: $\forall v \in L$ , reduce $v$ by $p$ 4: if any $v \in L$ is modified then 5: move $v$ to S 6: Loop <sup>†</sup> { $\forall v_1, v_2 \in L$ , reduce $p$ by $v_1, v_2$ } 7: if $p = 0$ then Return $p$ 8: $\forall v_1, v_2 \in L$ , reduce $v_1$ by $p, v_2$ 9: if any $v_1 \in L$ is modified then 10: move $v_1$ to S 11: Return $p$

<sup>&</sup>lt;sup>†</sup>After each pass of reducing p by all  $v \in L$  (respectively, by all pairs  $v_1, v_2 \in L$ ), we repeat the procedure as the current p may not be reduced with respect to some v (respectively, some pair of  $v_1, v_2$ ) in the list L any more. Note that the vector p is being updated from every reduction by v (respectively, pairs of  $v_1, v_2$ ). We repeat the loop (Lines 1 and 6 of Algorithm 6) until p can not be reduced any more with any  $v \in L$ (respectively, any pair  $v_1, v_2 \in L$ ).

Lines 1–5 of Algorithm 6 ensure the Gauss-reduced condition; lines 6–10 of Algorithm 6 ensure the 3-dimensional Minkowski reducedness. Note that the Triple MinkowskiSieve resembles the GaussSieve. In particular, we do not Minkowski reduce the three vectors in one go. Instead, we only modify one vector at each reduction step. If the modified vector comes from a new sampled vector p, then we repeatedly reduce p using every  $v_1, v_2 \in L$  (Line 6 of Algorithm 6) until it can not be reduced any more. If the modified vector comes from an existing list vector  $v \in L$ , then we move the reduced v' from the list L to the stack S for further consideration (Lines 9–10 of Algorithm 6).

During the execution of the Algorithm 5, every triple of vectors of L is always Minkowski reduced (except during the calls to Algorithm 6). We can terminate the algorithm if we have found a short enough vector or if the number of zero vectors returned by Algorithm 6) reaches some threshold (this determines the 'cond' in Line 2 of Algorithm 5).

As GaussSieve is one of the most promising lattice sieving candidates for use in practice, we implemented the triple and quadruple MinkowskiSieve algorithms. We also consider an improved variant of Triple MinkowskiSieve by applying the filtering principle (see §4) on pairs of vectors  $(\boldsymbol{p}, \boldsymbol{v}_1)$ . We refer to §6.2 for experimental results.

#### 6. Experiments

In this section, we describe some experimental results that support our previous analysis. In § 6.1, we conduct experiments to verify the conclusions in §§ 3.1 and 3.2. In § 6.2, we describe the implementation and experimental results for triple, quadruple and filtered triple MinkowskiSieve.

#### 6.1. Experiments on the unit sphere

We give some numerical evidence for the conclusions in § 3, in the case of list vectors sampled uniformly on the unit sphere. Note that, even in that idealized framework, the analysis of § 3 remains heuristic when  $k \ge 3$ .

We sample random vectors uniformly on the sphere and, for each new vector, we test whether it coincides with (for example, is close to) a combination of two vectors (respectively, one vector) already present in the list, in the case of triples (respectively, pairs).

We enforce Minkowski's reduction condition for the double- or triple-reduced vectors, and thus this is even stricter than the cost analysis. Once the number of 'collisions' (nearby vectors) exceeds a small multiple of  $|\mathbf{L}|$  (here we use  $4 \cdot |\mathbf{L}|$ ) we stop and record the list size. We repeat this process  $\lceil 20000/n \rfloor$  many times for each dimension n and take the average (logarithmic) list size and plot it in Figure 2.

The lines in the plot take the form  $C_1n + b (\log_2 n)^{e_1} + c$  and  $C_2n + d (\log_2 n)^{e_2} + e$ . We used a least-squares fit to find the best fits for parameters b, c, d, e. For the experimental data, the values  $C_1$  and  $C_2$  are 0.21 and 0.17, which are close to the heuristic estimates. It is also clear that there is a significant gap between pairwise and triplewise reduced list sizes.

#### 6.2. Implementation of the tuple MinkowskiSieve

We have implemented the GaussSieve, triple MinkowskiSieve, quadruple MinkowskiSieve and filtered triple MinkowskiSieve algorithms. These algorithms were described in § 5. The implementation<sup>†</sup> is based on FPLLL's codebase [4]. We use DGS [3] for sampling initial lattice vectors. We describe some experimental results for these algorithms.

 $<sup>^{\</sup>dagger}\mathrm{The}$  implementation will be incorporated into FPLLL in the next release.

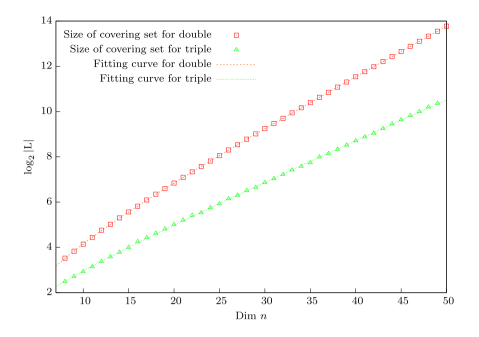


FIGURE 2. Log space complexity of double and triple sieve.

For each dimension, we use at least ten random instances from the SVP challenge generator [6]. Before the sieving, we pre-process the input basis with lattice reduction. To prevent trivial instances (where the sieving quickly finds many collisions, since the vectors are already short), we used either LLL with  $\delta = 0.75$ ,  $\eta = 0.51$  for smaller dimensions or BKZ (block Korkin–Zolotarev) with blocksize 20 for larger dimensions. The computations are taken on Intel Xeon X5650 processors of 2.67 GHz.

We give experimental results in Table A.1 and Figures A.1 and A.2 (see the appendix). We explain the notation in Table A.1 (Figures A.1 and A.2 follow a similar notation). The column n denotes the dimension. Column '2-red' denotes GaussSieve, Column '3-red' denotes triple MinkowskiSieve, '4-red' denotes quadruple MinkowskiSieve and Column '3-red' is the filtered triple MinkowskiSieve. Under each algorithm, the subcolumn  $|L|_{\infty}$  denotes the (average) maximum list size during the sieving. The number inside parentheses (for example, (.251) in n = 24 and 2-red) is the average value of  $(\log |L|_{\infty})/n$  over all experiments. Subcolumn time is the average running time in seconds. The cells with 'b' are the experiments with BKZ-20 pre-processed; other cells are LLL pre-processed.

It can be seen that there is a noticeable gap between the space requirements of GaussSieve and triple MinkowskiSieve (respectively, quadruple MinkowskiSieve). Furthermore, the filtered triple MinkowskiSieve variant is much more efficient than the non-filtered triple MinkowskiSieve.

# 7. Discussion

In the present work, we describe variant tuple sieving algorithms which reduce the memory requirement in lattice sieving. For triple (quadruple) reduced vectors, we estimate the space complexity to be  $2^{0.1887n+o(n)}$  (respectively,  $2^{0.1724n+o(n)}$ ). The (heuristic) asymptotic costs of these algorithms are investigated and verified in experiments.

One interesting future question is to consider nearest neighbor search techniques to speed up the search procedures, which could lead to an improvement on the overall running time of the sieving algorithms. For triple sieving, for instance, we could consider a nest of hash functions, where the first search is to find vectors with inner product at least 1/3 and the second search is to find vectors with inner product 1/2. The two sets of hash tables  $\mathcal{T}_{1/3}, \mathcal{T}_{1/2}$  are therefore different; one is optimized for finding vectors at angle  $\approx 70^{\circ}$  and one for 60°.

# Appendix

The appendix contains experimental results from  $\S 6.2$ .

	2-red		3-red		3-red'		4-red	
n	$ L _{\infty}$	Time	$ L _{\infty}$	Time	$ L _{\infty}$	Time	$ L _{\infty}$	Time
24	70(0.251)	0.02	42 (0.222)	0.11	50 (0.233)	0.03	33 (0.208)	1.1e1
26	89(0.238)	0.04	49 (0.209)	0.21	57 (0.217)	0.04	34 (0.190)	2.5e1
28	130(0.246)	0.05	74(0.222)	0.41	83 (0.227)	0.07	55 (0.206)	6.8e1
30	165 (0.238)	0.08	81 (0.207)	0.95	94 (0.214)	0.13	58 (0.192)	1.5e2
32	244 (0.248)	0.11	97~(0.200)	1.8	123 (0.213)	0.22	$62 \ (0.183)$	3.2e2
34	319(0.245)	0.19	148 (0.212)	4.8	169(0.217)	0.48	97~(0.194)	9.6e2
36	435(0.243)	0.31	176(0.207)	9.9	223 (0.217)	0.89	118 (0.191)	2.6e3
38	571(0.241)	0.61	$230 \ (0.206)$	$2.7\mathrm{e}1$	284 (0.214)	1.9	$134 \ (0.186)$	6.5e3
40	741 (0.238)	1.1	298 (0.205)	6.3e1	361 (0.212)	4.4	191 (0.189)	1.9e4
42	$1021 \ (0.238)$	2.3	377 (0.204)	1.6e2	476 (0.212)	9.4	ь 246 (0.189)	2.0e4
44	1390(0.237)	5.1	484 (0.203)	4.0e2	602 (0.219)	$2.3\mathrm{e1}$	ь 263 (0.182)	5.2e4
46	1777 (0.235)	9.6	638 (0.202)	9.9e2	777 (0.209)	$4.7\mathrm{e}1$	ь 352 (0.184)	1.4e5
48	2400 (0.234)	2.0e1	795~(0.201)	2.5e3	1063 (0.209)	1.1e2		—
50	3254(0.233)	4.1e1	1104 (0.202)	6.6e3	1328 (0.207)	2.5e2		
52	4219 (0.232)	9.5e1	$1324 \ (0.199)$	1.7e4	1742 (0.207)	5.6e2		—
54	5879(0.232)	2.2e2	$1700 \ (0.199)$	3.9e4	2234 (0.206)	1.2e3		—
56	7574(0.230)	5.3e2	ь 2163 (0.198)	2.7e4	ь 2878 (0.205)	8.5e2		
58	10539 (0.230)	1.2e3	ь 2877 (0.198)	7.1e4	ь 3804 (0.205)	1.9e3		—
60	ь 13433 (0.229)	9.1e2	ь 3664 (0.197)	1.7e5	ь 4879 (0.204)	4.3e3		
62	ь 18251 (0.228)	2.0e3	ь 5102 (0.199)	4.6e5	ь 6475 (0.204)	1.0e4		
64	ь 24223 (0.228)	4.5e3	ь 6604 (0.198)	1.1e6	ь 8338 (0.204)	2.2e4		—
66	ь 32587 (0.227)	9.1e3			ь 10994 (0.203)	$5.1\mathrm{e}4$		
68	ь 43887 (0.227)	2.0e4		_	ь 14297 (0.203)	1.1e5		
70	ь 58912 (0.227)	3.8e4		_	ь 18973 (0.203)	2.5e5		—
72	ь 79521 (0.226)	7.2e4		_				
74	ь 107050 (0.226)	1.5e5						
76	ь 142504 (0.225)	2.9e5						
78	ь 192141 (0.225)	6.0e5					—	
80	ь 256343 (0.225)	1.2e6						—

TABLE A.1. Experimental results for (filtered) tuple MinkowskiSieve ( $\S 6.2$ ).

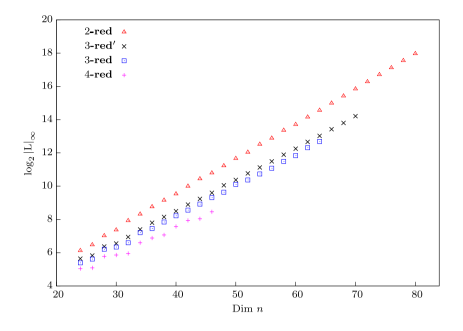


FIGURE A.1. Plot of logarithmic average maximum list size  $(\log_2 |L|_{\infty})$  for Table A.1.

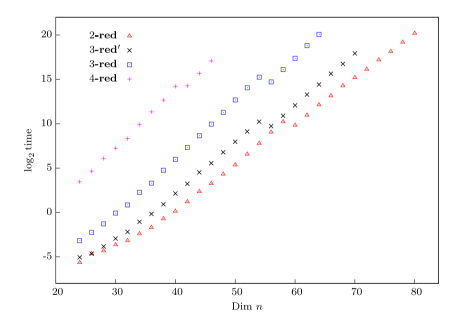


FIGURE A.2. Plot of logarithmic average time ( $\log_2 time$ ) for Table A.1. Note that we used either LLL or BKZ-20 for the pre-processing, depending on the dimension of the input instance.

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