A sieve algorithm based on overlattices

Anja Becker, Nicolas Gama and Antoine Joux

ABSTRACT

In this paper, we present a heuristic algorithm for solving exact, as well as approximate, shortest vector and closest vector problems on lattices. The algorithm can be seen as a modified sieving algorithm for which the vectors of the intermediate sets lie in overlattices or translated cosets of overlattices. The key idea is hence no longer to work with a single lattice but to move the problems around in a tower of related lattices. We initiate the algorithm by sampling very short vectors in an overlattice of the original lattice that admits a quasi-orthonormal basis and hence an efficient enumeration of vectors of bounded norm. Taking sums of vectors in the sample, we construct short vectors in the next lattice. Finally, we obtain solution vector(s) in the initial lattice as a sum of vectors of an overlattice. The complexity analysis relies on the Gaussian heuristic. This heuristic is backed by experiments in low and high dimensions that closely reflect these estimates when solving hard lattice problems in the average case.

This new approach allows us to solve not only shortest vector problems, but also closest vector problems, in lattices of dimension \( n \) in time \( 2^{0.8774 n} \) using memory \( 2^{0.2925 n} \). Moreover, the algorithm is straightforward to parallelize on most computer architectures.

1. Introduction

Hard lattice problems, such as the shortest vector problem (SVP) and the closest vector problem (CVP), have a long-standing relationship to number theory and cryptology. In number theory, they can for example be used to find Diophantine approximations. In cryptology, they were used as cryptanalytic tools for a long time, first through a direct approach as in [20] and then more indirectly using Coppersmith’s small roots algorithms [8, 9]. More recently, these hard problems have also been used to construct cryptosystems. Lattice-based cryptography is also a promising area due to the simple additive, parallelizable structure of a lattice. The two basic hard problems SVP and CVP are known to be NP-hard† to solve exactly [1, 22] and also NP-hard to approximate [10, 26] within at least constant factors. The time complexity of known algorithms that find the exact solution is at least exponential in the dimension of the lattice. These algorithms also serve as subroutines for strong polynomial time approximation algorithms. Algorithms for the exact problem hence enable us to choose appropriate parameters.

A shortest vector can be found by enumeration [21, 36], sieving [3, 28, 31, 38] or the Voronoi-cell algorithm [27]. Enumeration uses a negligible amount of memory and its running time is between \( n^{O(n)} \) and \( 2^{O(n^2)} \) depending on the amount and quality of the preprocessing. Probabilistic sieving algorithms, as well as the deterministic Voronoi-cell algorithm are simply exponential in time and memory. A closest vector can be found by enumeration and by the Voronoi-cell algorithm; however, to the best of our knowledge, no sieve algorithm is known

Received 27 February 2014; revised 23 May 2014.

2010 Mathematics Subject Classification 11H99 (primary), 11Y16, 68T20, 90C59 (secondary).

Contributed to the Algorithmic Number Theory Symposium XI, GyeongJu, Korea, 6–11 August 2014.

Part of this work was supported by the Swiss National Science Foundation under grant numbers 200021-126368 and 200020-153113.

†Under randomized reductions in the case of SVP.
to provably solve CVP instances, and it would be interesting to study other sieve algorithms which also work in the CVP case. Table 1 presents the complexities of currently known SVP and CVP algorithms including our new algorithm. In particular, it shows that the asymptotic time complexity of our new approach (slightly) outperforms the complexity of the best pre-existing sieving algorithm and that, as a bonus, it can for the same price serve as a CVP algorithm. The high memory requirement limits the size of accessible dimensions, for example we need 3 TB of storage in dimension 90 which we divide into 25 groups of 120 GB in RAM, and we would need twice as much in dimension 96. For this reason, the algorithm, as well as other classical sieving techniques, is in practice not competitive with the fastest memoryless methods such as pruned enumeration or aborted BKZ. However, our experiments suggest that despite the higher memory requirements, the sequential running time of our algorithm is of the same order of magnitude as the Gauss sieve, but with an algorithm easier to parallelize.

A long-standing open question was to find ways to decrease the complexity of enumeration-based algorithms to a single exponential time complexity. On an LLL- or BKZ-reduced basis [24, 36] the running time of Schnorr and Euchner’s enumeration is doubly exponential in the dimension. If we further reduce the basis to an HKZ-reduced basis [23], the complexity becomes $2^{O(n \log n)}$ [18, 21]. Enumeration would become simply exponential if a quasi-orthonormal basis, as defined in §2, could be found. Unfortunately, most lattices do not possess such a favorable quasi-orthonormal basis. Also for random lattices the lower bound on the Rankin invariant is of size $2^{O(n \log n)}$ and determines the minimal complexity for enumeration that operates exclusively on the original lattice. We provide a more detailed discussion in §2.

Our approach circumvents this problem by making use of overlattices that admit a quasi-orthonormal basis. These overlattices are found in polynomial time by an algorithm relying on the structural reduction introduced by [14], as described in §3.2. Once we have an overlattice and its quasi-orthonormal basis, we may efficiently enumerate short vectors at a constant factor of the first minimum in the overlattice. Our main task is to turn these small samples into a solution vector in the initial lattice. The construction is very similar to an observation by Mordell [30] in 1935 which presented the first algorithmic proof of Minkowski’s inequality using only finite elements. Namely, he observed that given a lattice $\mathcal{L}_i$ and an overlattice $\mathcal{L}_{i+1} \supset \mathcal{L}_i$ such that $[\mathcal{L}_{i+1} : \mathcal{L}_i] = r$, in any pool of at least $r + 1$ short vectors of $\mathcal{L}_{i+1}$, there exist at least two vectors whose difference is a short non-zero vector in $\mathcal{L}_i$. This construction has also been implicitly used in worst-case to average-case reductions, where a short overlattice basis is used to sample a pool of short Gaussian overlattice vectors, which are then combined by an SIS (short integer solution) oracle into polynomially longer vectors of the original lattice. In our setting, the overlattice basis is quasi-orthonormal, which allows an efficient enumeration

### Table 1. Complexity of currently known SVP/CVP algorithms.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Time</th>
<th>Memory</th>
<th>CVP</th>
<th>SVP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kannan enumeration [18]</td>
<td>$n^{n/2+o(n)}$</td>
<td>$\text{poly}(n)$</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td></td>
<td>$n^{n/(2e)+o(n)}$</td>
<td>$\text{poly}(n)$</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Voronoï cell [27]</td>
<td>$2^{2n}$</td>
<td>$2^n$</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>ListSieve birthday [33]</td>
<td>$2^{2.465 n + o(n)}$</td>
<td>$2^{1.233 n + o(n)}$</td>
<td>×</td>
<td>✓</td>
</tr>
<tr>
<td>GaussSieve [28]</td>
<td>$2^{0.415 n + o(n)}$</td>
<td>$2^{0.2075 n + o(n)}$</td>
<td>?</td>
<td>✓</td>
</tr>
<tr>
<td>Nguyen–Vidick sieve [31]</td>
<td>$2^{0.415 n + o(n)}$</td>
<td>$2^{0.2075 n + o(n)}$</td>
<td>×</td>
<td>✓</td>
</tr>
<tr>
<td>WLTB sieve [38]</td>
<td>$2^{0.3836 n + o(n)}$</td>
<td>$2^{0.2557 n + o(n)}$</td>
<td>×</td>
<td>✓</td>
</tr>
<tr>
<td>Three-level sieve [39]</td>
<td>$2^{0.3778 n + o(n)}$</td>
<td>$2^{0.2833 n + o(n)}$</td>
<td>×</td>
<td>✓</td>
</tr>
<tr>
<td>Our algorithm</td>
<td>from $2^{0.4150 n}$ to $2^{0.3774 n}$</td>
<td>$2^{0.2075 n}$</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

---

---

Downloaded from https://www.cambridge.org/core. IP address: 54.70.40.11, on 28 Jun 2019 at 19:07:43, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1112/S1461157014000029
of the shortest overlattice vectors. These vectors are then combined to the shortest vectors of the original lattice by a concrete, albeit exponential time, algorithm.

The new algorithm solves SVP and CVP for random lattices in the spirit of a sieving algorithm, except that intermediate vectors lie in overlattices or cosets of overlattices whose geometry varies from dense lattices to quasi-orthogonal lattices. Alternatively, the algorithm can be viewed as an adaptation of the representation technique that solves knapsack problems \[4\] and decoding problems \[5, 25\] to the domain of lattices. Due to the richer structure of lattices, the adaptation is far from straightforward. To give a brief analogy, instead of searching for a knapsack solution, assume that we want to find a short vector in an integer lattice. An upper bound on the Euclidean norm of the solution vector provides a geometric constraint, which induces a very large search space. The short vector we seek can be decomposed in many ways as the sum of two shorter vectors with integer coefficients. Assuming that these sums provide \(N\) different representations of the same solution vector, we can then choose any arbitrary constraint which eliminates all but a fraction \(\approx 1/N\) of all representations. With this additional constraint, the solution vector can still be efficiently found, in a search space reduced by a factor \(N\). From a broader perspective, this technique can be used to transform a problem with a hard geometric constraint, like short lattice vectors, into an easier subproblem, like short integer vectors (because \(\mathbb{Z}^n\) has an orthonormal basis), together with a custom additional constraint which is in general linear or modular, allowing an efficient recombination of the solutions to the subproblems. The biggest challenge is to bootstrap the algorithm by finding suitable and easier subproblems using overlattices. We propose a method that achieves this thanks to a well-chosen overlattice allowing an efficient deterministic enumeration of vectors of bounded norm. In this way, we can compute a starting set of vectors that is used to initiate a sequence of recombinations that ends up solving the problem initially considered.

**Our contribution.** We present a new heuristic algorithm for the exact SVP and CVP for \(n\)-dimensional lattices using a tower of \(k\) overlattices \(L_i\), where \(L = L_0 \subseteq \ldots \subseteq L_k\). In this tower, we choose the lattice \(L_k\) at the bottom of the tower in a way that ensures that we can efficiently compute a sufficiently large pool of very short vectors in \(L_k\). Starting from this pool of short vectors, we move from each lattice of our tower to the one above using summation of vectors while controlling the growth of norms.

For random lattices and under heuristic assumptions, two \(L_{i+1}\)-vectors sum up to an \(L_i\)-vector with probability \(1/\alpha^n\), where \(\text{vol}(L_i)/\text{vol}(L_{i+1}) = \alpha^n > 1\). We allow the norm to increase by a moderate factor \(\alpha\) in each step, in order to preserve the size of our pool of available vectors per lattice in our tower.

Our method can be used to find vectors of bounded norm in a lattice \(L\) or, alternatively, in a coset \(x + L\), \(x \notin L\). Thus, in contrast to classical sieving techniques, it allows us to solve both SVP or CVP, and more generally, to enumerate all lattice points within a ball of fixed radius. The average running time in the asymptotic case is \(2^{0.3774n}\), requiring a memory of \(2^{0.2925n}\). It is also possible to choose different time–memory tradeoffs and devise slower algorithms that need less memory. We report our experiments on random lattices and SVP challenges of dimension 40 to 90, whose results confirm our theoretical analysis and show that the algorithm works well in practice. We also study various options to parallelize the algorithm and show that parallelization works well on a wide range of computer architectures.

2. **Background and notation**

**Lattices and cosets.** A lattice \(L\) of dimension \(n\) is a discrete subgroup of \(\mathbb{R}^m\) that spans an \(n\)-dimensional linear subspace. A lattice can be described as the set of all integer combinations
\{\sum_{i=1}^{n} \alpha_i b_i \mid \alpha_i \in \mathbb{Z}\} \text{ of } n \text{ linearly independent vectors } b_i \text{ of } \mathbb{R}^m. \text{ Such vectors } b_1, \ldots, b_n \text{ are called a basis of } \mathcal{L}. \text{ The volume of the lattice } \mathcal{L} \text{ is the volume of } \text{span}(\mathcal{L})/\mathcal{L}, \text{ and can be computed as } \sqrt{\det(BB^t)}, \text{ for any basis } B. \text{ Any lattice has a shortest non-zero vector of Euclidean length } \lambda_1(\mathcal{L}) \text{ which can be upper-bounded by Minkowski's theorem as } \lambda_1(\mathcal{L}) \leq \sqrt{n} \cdot \text{vol}(\mathcal{L})^{1/n}. \text{ We call a coset of } \mathcal{L} \text{ a translation } x + \mathcal{L} = \{x + v \mid v \in \mathcal{L}\} \text{ of } \mathcal{L} \text{ by a vector } x \in \text{span}(\mathcal{L}).

**Overlattice and index.** A lattice \( \mathcal{L}' \) of dimension \( n \) such that \( \mathcal{L} \subseteq \mathcal{L}' \) is called an overlattice of \( \mathcal{L} \). The quotient group \( \mathcal{L}' / \mathcal{L} \) is a finite abelian group of order \( \text{vol}(\mathcal{L})/\text{vol}(\mathcal{L}') = [\mathcal{L}' : \mathcal{L}] \).

**Hyperballs.** Let \( \text{Ball}_n(R) \) denote the ball of radius \( R \) in dimension \( n \), where we omit \( n \) if it is implied by the context. The volume \( V_n \) of the \( n \)-dimensional ball of radius 1 and the radius \( r_n \) of the \( n \)-dimensional ball of volume 1 are respectively:

\[
V_n = \frac{\sqrt{\pi^n}}{\Gamma(n/2 + 1)} \quad \text{and} \quad r_n = V_n^{-1/n} = \sqrt{n/2\pi^e(1 + o(1))}.
\]

**Gaussian heuristic.** In many cases, when we wish to estimate the number of lattice points in a ‘nice enough’ set \( S \), we can use the Gaussian heuristic. In this paper, we will quantify the Gaussian heuristic as follows:

**HEURISTIC 2.1 (Gaussian heuristic).** There exists a constant \( G_H \geq 1 \) such that for all the lattices \( \mathcal{L} \) and all the sets \( S \) that we consider in this paper, the number of points in \( S \cap \mathcal{L} \) satisfies:

\[
\frac{1}{G_H} \cdot \frac{\text{vol}(S)}{\text{vol}(\mathcal{L})} \leq \#(S \cap \mathcal{L}) \leq G_H \cdot \frac{\text{vol}(S)}{\text{vol}(\mathcal{L})}.
\]

In fact, it can be proved that for any bounded measurable set \( S \), the expectation, over random unit volume lattices drawn from the Haar distribution \([2]\), of the number of non-zero points is always the volume of \( S \). Also, for any fixed lattice of unit volume and any fixed bounded measurable set \( S \), the expectation of the number of lattice points in \( t + S \) for uniform \( t \) modulo \( \mathcal{L} \) is exactly \( \text{vol}(S) \). However, fewer results are known about the standard deviation, or whether these distributions are concentrated enough around the expectation so that almost all instances satisfy the upper and lower bounds.

In this paper, the geometry of lattices varies between random integer lattices of large enough volume and quasi-orthonormal lattices. We will assume that in these lattices, the length \( \lambda_1 \) of the shortest vector is the one given by the Gaussian heuristic, that is, the radius of a ball of volume \( \text{vol}(\mathcal{L}) \): \( \lambda_1(\mathcal{L}) \approx r_n \cdot \sqrt{\text{vol}(\mathcal{L})} \). Furthermore, the sets \( S \) we consider in this paper are either balls of radius larger than \( \sqrt{3/2} r_n \cdot \sqrt{\text{vol}(\mathcal{L})} \), and whose center is uniform modulo the lattice (that is, far from \( 0 \)), or the intersection of two such balls whose centers are close enough. In these cases, our experiments indicate that the number of lattice points in these sets is almost always between 50% and 110% of \( \text{vol}(S)/\text{vol}(\mathcal{L}) \), thus Heuristic 2.1 holds in practice for \( G_H = 2 \).

**Gram–Schmidt orthogonalization (GSO).** The GSO of a non-singular square matrix \( B \) is the unique decomposition as \( B = \mu \cdot B^* \), where \( \mu \) is a lower triangular matrix with unit diagonal and \( B^* \) consists of mutually orthogonal rows. For each \( i \in \{1, \ldots, n\} \), we call \( \pi_i \) the orthogonal projection over \( \text{span}(b_1, \ldots, b_{i-1}) \). In particular, one has \( \pi_i(b_i) = b_i^* \), which is the \( i \)th row of \( B^* \). We use the notation \( B_{[i,j]} \) for the projected block \( [\pi_i(b_i), \ldots, \pi_i(b_j)] \).

\( ^\dagger \)The algorithms and proofs would also work with \( G_H = \text{poly}(n) \) or \( G_H = (1 + \varepsilon)^n \), giving slightly worse complexities.
Rankin factor and quasi-orthonormal basis. Let \( B \) be an \( n \)-dimensional basis of a lattice \( \mathcal{L} \), and \( j \leq n \). We call the ratio
\[
\gamma_{n,j}(B) = \frac{\operatorname{vol}(B_{[1,j]})}{\operatorname{vol}(\mathcal{L})^{j/n}} = \frac{\operatorname{vol}(\mathcal{L})^{(n-j)/n}}{\operatorname{vol}(\pi_{j+1}(\mathcal{L}))}
\]
the Rankin factor of \( B \) with index \( j \). The well-known Rankin invariants of the lattice, \( \gamma_{n,j}(\mathcal{L}) \), introduced by Rankin [34], are simply the squares of the minimal Rankin factors of index \( j \) over all bases of \( \mathcal{L} \). This allows us to define a quasi-orthonormal basis.

**Definition 2.2 (Quasi-orthonormal basis).** A basis \( B \) is **quasi-orthonormal** if and only if its Rankin factors satisfy \( 1 \leq \gamma_{n,j}(B) \leq n \) for all \( j \in [1, n] \).

In the above definition, we chose the upper bound \( n \) over a general \( \poly(n) \) only because we are able to achieve this factor. More generally, any polynomial function would be sufficient for the asymptotical analysis and for the running time. For example, any real triangular matrix with identical diagonal coefficients forms a quasi-orthogonal basis. More generally, any basis whose \( \|b_i^*\| \) are almost equal is quasi-orthogonal. This is a very strong notion of reduction, since average LLL-reduced or BKZ-reduced bases only achieve a \( 2^{O(n^2)} \) Rankin factor and HKZ-reduced bases of random lattices have a \( 2^{O(n \log n)} \) Rankin factor. Finally, Rankin's invariants are lower-bounded [6, 13, 37] by \( 2^{\Theta(n \log n)} \) for almost all lattices, which means that only lattices in a tiny subclass possess a quasi-orthonormal basis.

**Schnorr–Euchner enumeration.** Given a basis \( B \) of an integer lattice \( \mathcal{L} \subseteq \mathbb{R}^n \), the Schnorr–Euchner enumeration algorithm [36] allows us to enumerate all vectors of Euclidean norm \( \leq R \) in the bounded coset \( C = (z + \mathcal{L}) \cap \text{Ball}_R(R) \) where \( z \in \mathbb{R}^n \). The running time of this algorithm is
\[
T_{\text{SE}} = \sum_{i=1}^{n} \#((\pi_{n+1-i}(z + \mathcal{L}) \cap \text{Ball}_R(R))
\]
which is equivalent to
\[
T_{\text{SE}} \approx \sum_{i=1}^{n} \frac{\operatorname{vol}(\text{Ball}_R(R))}{\operatorname{vol}(\pi_{n+1-i}(\mathcal{L}))}
\]
under Heuristic 2.1. The last term in the sums (2.1) and (2.2) denotes the number of solutions \( \#C \). Thus, the complexity of enumeration is approximately \( T_{\text{SE}} \approx \tilde{O}(\#C) \cdot \max_{j \in [1,n]} \gamma_{n,j}(B) \).

This is why a reduced basis of smallest Rankin factor is favorable. The lower bound on Rankin's invariant of \( \gamma_{n,n/2}(\mathcal{L}) = 2^{\Theta(n \log n)} \) for most lattices therefore determines the minimal complexity of enumeration that is achievable while working with the original lattice, provided that one can actually compute a basis of \( \mathcal{L} \) minimizing the Rankin factors, which is also NP-hard. If the input basis is quasi-orthonormal, the upper bound \( \gamma_{n,j}(B) \leq n \) from Definition 2.2 implies that the enumeration algorithm runs in time \( \tilde{O}(\#C) \), which is optimal. Without knowledge of a good basis one can aim to decompose the problem into more favorable cases that finally allow us to apply the Schnorr–Euchner algorithm as we describe below.

3. **Enumeration of short vectors by intersection of hyperballs**

This section presents the new algorithm that enumerates \( \beta^n \) shortest vectors in any coset \( t + \mathcal{L} \) of a lattice \( \mathcal{L} \) for a constant \( \beta \approx \sqrt{3/2} \). It can be used to solve the NP-hard problems
\[^{\dagger}\gamma_{2n,n}(\mathcal{L}) \geq (n/12)^n \text{ with probability } \approx 1 \text{ on random real lattices of volume } 1 \text{ drawn from the Haar distribution.}\]
SVP, CVP, ApproxEVPβ 3 and ApproxCVPβ: given a lattice L, the SVP can be reduced to enumerating vectors of Euclidean norm O(λ 1(L)) in the coset 0 + L while a CVP instance can be solved by enumerating vectors of norm at most dist(t, L) in the coset −t + L. These bounded cosets, (t + L) ∩ Balln(R) for suitable radius R, can be constructed in an iterative way by use of overlattices. The searched vectors are expressed as a sum of short vectors of suitable translated overlattices of smaller volume. The search for a unique element in a lattice as required in the SVP or CVP is delegated to the problem of enumerating bounded cosets. Any non-trivial element found by our algorithm is naturally a solution to the corresponding ApproxEVPβ or ApproxCVPβ.

We present the new algorithm solving lattice problems based on intersections of hyperballs in §3.1 and the generic initialization of our algorithm as described in §3.2. Finally, §3.3 describes the cost of the first step in the algorithm.

3.1. General description of the new algorithm

Assume that we are given a tower of k = O(n) lattices Li ⊂ Rn of dimension n where Li ⊆ Li+1 and the volume of any two consecutive lattices differs by a factor N = [αn] ∈ N>1. We also assume that the bottom lattice Lk permits an efficient enumeration of the βn shortest vectors in any coset t + Lk for 1 < β < √3/2. The ultimate goal is to find the βn shortest vectors in some coset t0 + L0 of L0. We postpone how to find suitable lattices Li, i ≥ 1, to the following two sections.

We also assume in this section, that the Gaussian heuristic (Heuristic 2.1) holds. Under this assumption, the problem of finding the βn shortest elements in some coset t + L is roughly equivalent to enumerating all lattice vectors of L in the ball of radius β · r n · √vol(L) centered at −t ∈ Rn.

Each step for i = k − 1 down to 0 of the algorithm is based on the following intuition: we are given the βn shortest vectors v j in ti/2 + Li+1. By summation, we can then find vectors (v j + v i) j∈I that lie in ti + Li+1. We select those which actually lie in ti + Li and have norm small enough, and consider them as the input pool for the next step. For suitable parameters, namely α small enough and β large enough, we thus recover the βn shortest vectors of ti + Li.

More precisely, for each i ∈ [0, k], we call Ci the bounded coset of Li that contains the βn shortest vectors of the coset ti + Li where ti = t0/2i ∈ Rn. More formally, let us define

\[ R_i = β · r_n · √vol(L_i) \quad \text{and} \quad C_i = (t_i + L_i) ∩ Ball(0, R_i) \]

such that

\[ #C_i ≈ vol(Ball(R_i))/vol(L_i) = β^n, \]

which follows from Heuristic 2.1. In addition, we recall that

\[ L_i ⊂ L_{i+1} \quad \text{where} \quad vol(L_i)/vol(L_{i+1}) = [α^n]. \]

In order to enumerate C0, our algorithm successively enumerates C i, starting from i = k down to zero; Figure 1 illustrates the sequence of enumerated lists.

During the construction of the tower of lattices, which is studied in the next sections, we already ensure that Ck is easy to obtain. We now explain how we can compute Ci−1 from Ci. To do this, we compute all sums x + y of vector pairs of Ci × Ci which satisfy
A sieve algorithm based on overlattices

\[ C_0 \]
\[ + \text{ check (3.1), (3.2)} \]
\[ C_i \]
\[ + \text{ check (3.1), (3.2)} \]

Enumerate \( C_k \)

**Figure 1. Iterative creation of lists.**

the conditions

\[ x + y \in t_{i-1} + L_{i-1} \text{ and } \parallel x + y \parallel \leq \beta \cdot r_n \cdot \sqrt{\text{vol}(L_{i-1})}. \] (3.2)

This means that we collect the \( \beta^n \) shortest vectors of the coset \( C_{i-1} = t_{i-1} + L_{i-1} \) by going through \( C_i = t_i + L_i \). In practice, an equivalent way to check if condition (3.1) holds, is to use an efficient computation for the map \( \varphi_{i-1} : C_i \rightarrow L_i/L_{i-1}, z \rightarrow z - t_i \mod L_{i-1} \) and to verify that \( \varphi_{i-1}(x) + \varphi_{i-1}(y) = 0 \). Algorithm 1 summarizes our approach.

**Algorithm 1** Coset enumeration

| Constants: | \( \alpha \approx \sqrt{4/3}, \beta \approx \sqrt{3/2} \) |
| Parameters: | \( k \) |
| **Input:** | An LLL-reduced basis \( B \) of \( L_0 \) and a center \( t \in \mathbb{R}^n \) |
| **Output:** | Elements of \( t + L_0 \) of norm \( \leq R_0 = \beta r_n \text{vol}(L_0)^{1/n} \) |
| 1: | Randomize the input target by sampling \( t_0 \in t + L \). Use, for example, a discrete Gaussian distribution of parameter \( \sqrt{n} \| B^* \| \). This defines all the sub-targets \( t_i = t_0/2^i \) |
| 2: | Compute a tower of lattices \( L_0, \ldots, L_k \) by use of Algorithm 3 such that |
| | - \( L_0 \subset L_1 \subset \ldots \subset L_k \) and \( \text{vol}(L_i)/\text{vol}(L_{i-1}) = N = [\alpha^n] \) |
| | - lattice enumeration is easy on \( L_k \) |
| | - testing morphisms \( \varphi_{i-1} \) from \( t_i + L_i \) to \( L_i/L_{i-1} \) are efficient to evaluate. |
| 3: | Enumerate bottom coset \( C_k \) (with Schnorr–Euchner) |
| 4: | for \( i = k - 1 \) downto 0 do |
| 5: | \( C_i \leftarrow \text{Merge}(C_{i+1}, \varphi_i, R_i = \beta r_n \text{vol}(L_i)^{1/n}) \) (Algorithm 2) |
| 6: | end for |
| 7: | return \( C_0 \) |

A naive implementation of the merge routine that creates \( C_{i-1} \) from \( C_i \) would just run through the \( \beta^{2n} \) pairs of vectors from \( C_i \times C_i \), and eliminate those that do not satisfy the constraints (3.1) and (3.2). By regrouping the elements of \( C_i \) into \( \alpha^n \) buckets, according to their value modulo \( L_{i-1} \), condition (3.1) implies that each element of \( C_i \) only needs to be paired with the elements of a single bucket (see Algorithm 2). Heuristic 2.1 implies that each bucket contains at most \( G_H(\beta/\alpha)^n \) elements, therefore the merge operation can then be performed in time \( G_H^2(\beta^2/\alpha)^n \).

**Complexity and constraints for parameters \( \alpha \) and \( \beta \).** We now prove the complexity and correctness of Algorithm 2.
We thus require that $\operatorname{vol}(L) \in (0, \infty), \quad \text{and the second one in } C$, and output:

$$\sqrt{\frac{1}{2} - \frac{\alpha}{2}^2} \cdot \frac{1}{\sqrt{n}} \quad \text{ or alternatively } \quad \beta \sqrt{1 - \frac{\alpha}{2}^2} \geq 1 + \varepsilon_n \quad \text{ for } \varepsilon_n = (G_H \sqrt{n}/0.692)^{1/n} - 1 \text{ decreases towards } 0 \text{ as } n \text{ grows.}$$

Of course, for optimization reasons, we want to minimize the size of the lists $\beta^n$, and the number of steps $(\beta^2/\alpha)^n$ in the merge. Therefore we want to minimize $\beta$ and maximize $\alpha$ under the above constraint. The total running time of Algorithm 1 is given by $B + \operatorname{poly}(n)(\beta^2/\alpha)^n$ where $B$ represents the running time of the initial enumeration at level $k$ (details in §3.3). For optimal parameters, inequality (3.4) is in fact an equality. Asymptotically, the shortest running time occurs for $\alpha = \sqrt{\frac{4}{3}}$ and $\beta = \sqrt{\frac{3}{2}}$ for which a merge costs around $(\beta^2/\alpha)^n \approx 2^{0.3774 n}$ and the size of the lists is $\beta^n \approx 2^{0.2925 n}$. 

**Algorithm 2 Merge by collision**

```plaintext
/* Efficiently find pairs of vectors of $C_{i+1}$ s.t. their sum is in $C_i$ */
/* $C_i$ denotes $(t_i + L_i) \cap \operatorname{Ball}(R_i) */

\textbf{Input:} The bounded coset $C_{i+1}$, a testing morphism $\varphi_i$ and a radius $R_i$

\textbf{Output:} The bounded coset $C_i$

1: $C_i \leftarrow \emptyset$
2: Reorganize $C_{i+1}$ into buckets indexed by the values of $\varphi_i$
3: \textbf{for each } $v \in C_{i+1}$ \textbf{do}
4: \hspace{1em} \textbf{for each } $u$ in the bucket of index $-\varphi_i(v)$ \textbf{do}
5: \hspace{2em} \textbf{if } $\|u + v\| \leq R_i$ \textbf{then}
6: \hspace{3em} $C_i \leftarrow C_i \cup \{u + v\}$
7: \hspace{1em} \textbf{end if}
8: \textbf{end for}
9: \textbf{end for}
10: \textbf{return } C_i
```

**Theorem 3.1.** Assuming Heuristic 2.1, and provided that $\beta^n \geq G_H \sqrt{n}/0.692\sqrt{1 - \alpha^2/4^n}$, then, given as input the bounded coset $C_{i+1}$, Algorithm 2 outputs the coset $C_i$ within $G_H^2(\beta^2/\alpha)^n$ Euclidean norm computations. The memory is bounded by the size of the input and output: $G_H^2 \beta^n$ $n$-dimensional vectors.

**Proof.** It is clear that at each level, conditions (3.1) and (3.2) imply that Algorithm 2 outputs a subset of $C_i$. If we now need to prove that there exist constants $\alpha$ and $\beta$ such that all points of $C_i$ are present in the output. Equivalently, all points of $C_i$ must be expressed as the sum of two points in $C_{i+1}$ (see Figure 2 for an illustration). This geometric constraint can be simply rephrased as follows: a vector $z \in C_i$ is found if and only if there exists at least one vector $x$ of the coset $t_{i+1} + L_{i+1}$ in the intersection of two balls of radius $R_{i+1}$, the first one centered in 0, and the second one in $z$. It is clear that $z - x \in C_{i+1} = t_{i+1} + L_{i+1}$ since $2t_{i+1} = t_i$ and $L_i \subseteq L_{i+1}$. So if there is a point $x \in C_{i+1}$ in the intersection $I = \operatorname{Ball}(0, R_{i+1}) \cap \operatorname{Ball}(z, R_{i+1})$, we obtain $z \in C_i$ as a sum between $x \in C_{i+1}$ and $z - x \in C_{i+1}$. Under Heuristic 2.1, this occurs as soon as the intersection $I$ of the two balls has a volume larger than $G_H \operatorname{vol}(L_{i+1})$. We thus require that $\operatorname{vol}(I)/\operatorname{vol}(L_{i+1}) \geq G_H$. 

From Lemma A.1 and its corollary in the Appendix, we derive that the intersection of two balls of radius $R_i$ at distance at most $R_{i-1} = \alpha R_i$ is larger than $0.692 \cdot \operatorname{vol}(\operatorname{Ball}(R_i \cdot \sqrt{1 - (\alpha/2)^2}))/\sqrt{n}$. A sufficient condition on $\alpha$ and $\beta$ is then

$$\beta \sqrt{1 - (\alpha/2)^2} \geq (1 + \varepsilon_n) \quad \text{ for } \varepsilon_n = (G_H \sqrt{n}/0.692)^{1/n} - 1 \quad \text{ decreases towards } 0 \text{ as } n \text{ grows.}$$

Of course, for optimization reasons, we want to minimize the size of the lists $\beta^n$, and the number of steps $(\beta^2/\alpha)^n$ in the merge. Therefore we want to minimize $\beta$ and maximize $\alpha$ under the above constraint. The total running time of Algorithm 1 is given by $B + \operatorname{poly}(n)(\beta^2/\alpha)^n$ where $B$ represents the running time of the initial enumeration at level $k$ (details in §3.3). For optimal parameters, inequality (3.4) is in fact an equality. Asymptotically, the shortest running time occurs for $\alpha = \sqrt{\frac{4}{3}}$ and $\beta = \sqrt{\frac{3}{2}}$ for which a merge costs around $(\beta^2/\alpha)^n \approx 2^{0.3774 n}$ and the size of the lists is $\beta^n \approx 2^{0.2925 n}$. 

Downloaded from https://www.cambridge.org/core. IP address: 54.70.40.11, on 28 Jun 2019 at 19:07:43, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1112/S1461157014000229
A sieve algorithm based on overlattices

For any integer lattice $\mathcal{L}$, the simplest choice of an overlattice is $\mathcal{L}' = \mathbb{Z}^n$. However, this creates two problems.

1. $\mathbb{Z}^n$ does not satisfy our quantified version of Heuristic 2.1, as it would require an $n$-exponential $G_H$.

2. The index $[\mathbb{Z}^n : \mathcal{L}]$, which is the volume of $\mathcal{L}$, might be too large. Indeed, our sieve requires that the index is simply exponential, but an integer basis with polynomial entries may have a superexponential volume $2^{\text{poly}(n)}$.

For the first problem, it suffices to replace the square lattice $\mathbb{Z}^n$ by a quasi-orthogonal lattice $\mathcal{L}'$. Indeed, although the expectation of the number of lattice points in a randomly

Time–memory tradeoff. Other choices of $\alpha$ and $\beta$ that satisfy (3.4) provide a tradeoff between running time and required memory. Figure 3 shows the logarithmic size of the lists the algorithm needs to store depending on the time one is willing to spend. If one has access to only $\beta^n \approx 2^{0.21n}$ in memory, the time complexity increases to $(\beta^2/\alpha)^n \approx 2^{0.41n}$. In practice, we choose $\alpha > 1$ and $\beta > 0$ satisfying (3.3) with the constraint that $\alpha^n$ is integer.

3.2. Generic creation of the tower

For any integer $\mathcal{L}$, the simplest choice of an overlattice is $\mathcal{L}' = \mathbb{Z}^n$. However, this creates two problems.

1. $\mathbb{Z}^n$ does not satisfy our quantified version of Heuristic 2.1, as it would require an $n$-exponential $G_H$.

2. The index $[\mathbb{Z}^n : \mathcal{L}]$, which is the volume of $\mathcal{L}$, might be too large. Indeed, our sieve requires that the index is simply exponential, but an integer basis with polynomial entries may have a superexponential volume $2^{\text{poly}(n)}$.

For the first problem, it suffices to replace the square lattice $\mathbb{Z}^n$ by a quasi-orthogonal lattice $\mathcal{L}'$. Indeed, although the expectation of the number of lattice points in a randomly

Time vs. memory

Figure 3. Tradeoff between memory and time for varying choices of $\alpha$ and $\beta$.

Figure 2. Vector $z \in C_{i-1}$ found as sum between $x \in C_i$ and $z - x \in C_i \iff I \cap (t_i + \mathcal{L}_i) \neq \emptyset$. 
centered ball of radius $r_n/\beta \sqrt[1/n]{\text{vol}(\mathcal{L})}$ is always $\beta^n$, their repartition is much more concentrated around the expectation when the lattice is quasi-orthonormal than when it is $\mathbb{Z}^n$. In practice, almost all of these balls contain at least $\beta^n/2$ lattice points, and Heuristic 2.1 is valid with $G_H = 2$. The most noticeable exception is when the center is too close to a lattice point. In this very rare case, the number of points would exceed the upper bound by an exponential factor. Luckily, the lower bound in Heuristic 2.1 is more important than the upper bound: the lower bound is used an exponential number of times in equation (3.3) to prove the correctness of the merge, whereas the upper bound is used only with a polynomial number of different centers, to obtain the time and memory complexities. Thus bad centers are easy to avoid.

For the second problem, one can see that the difficulty of creating the sieve is not just to find a quasi-orthonormal overlattice (which is always possible), but to find one of small index. That is why we need a several overlattices instead of just one.

We now present a generic method of computing the tower of $\mathcal{L}_k$s that works well in practice for high dimensions, as we have verified in our experiments. Algorithm 3 summarizes the following steps.

We take as input a randomized LLL-reduced or BKZ-30-reduced basis $B$ of an $n$-dimensional lattice $\mathcal{L}$. We choose constants $\alpha > 1$ and $\beta > 0$ satisfying equation (3.4) with the additional constraint that $N = \alpha^n$ is an integer.

The Gram–Schmidt coefficients of $B$ usually decrease geometrically, and we can safely assume that $\min_i \|b_i^*\| \geq \max \|b_i^*\|/\sqrt{4/3}$. Otherwise, the LLL-reduced basis would immediately reveal a sublattice of dimension $< n$ containing the shortest vectors of $\mathcal{L}$. This means that there exists a smallest integer $k = \mathcal{O}(n)$ such that $\min_{i\in[1,n]} \|b_i^*\| \geq (\text{vol}(\mathcal{L})/N^k)^{1/n} = \sigma$. The integer $k$ determines the number of levels in our tower and $\sigma$ is the $n$th root of the volume of the last overlattice $\mathcal{L}_k$.

**Algorithm 3** Compute the tower of overlattices

**Input:** $B$, a (randomized) LLL-reduced basis of $\mathcal{L}$ of dimension $n$

**Output:** Bases $B^{(i)}$ of a tower of overlattices $\mathcal{L} = \mathcal{L}_0 \subset \ldots \subset \mathcal{L}_k$. Note that given a target $t_{i+1}$, the testing morphism $\varphi_i$ from $t_{i+1} + \mathcal{L}_{i+1}$ to $\mathbb{Z}_N$ is implicitly defined by $\varphi_i(t_{i+1} + \sum_{j=1}^n \mu_j b_j^{(i+1)}) = \mu_1 \mod N$.

1. Let $N = [\alpha^n]$.
2. Let $k$ be the smallest integer s.t. $N^k \geq \text{vol}(\mathcal{L})/\min_i \|b_i^*\|^n$.
3. Let $\sigma = (\text{vol}(\mathcal{L})/N^k)^{1/n}$, thus $\sigma \leq \min_i \|b_i^*\|$.
4. Apply Algorithm B.1 on input $(B, \sigma)$ to find a basis $\hat{B} = [\hat{b}_1, \hat{b}_2, \ldots, \hat{b}_n]$ of $\mathcal{L}$.
5. $B^{(i)} \leftarrow [\hat{b}_1/N^i, \hat{b}_2, \ldots, \hat{b}_n]$ for each $i \in [0, k]$.
6. return $B^{(i)}$ for all $i$.

It remains to find the tower $(\mathcal{L}_i)_{i\in[1,k]}$ of overlattices of $\mathcal{L}_i$ together with a quasi-orthonormal basis $B^{(k)}$ of $\mathcal{L}_k$, given a structural condition $\mathcal{L}^{(i)}/\mathcal{L}^{(i−1)} \simeq \mathbb{Z}/N \mathbb{Z}$ (Algorithm 3). This problem is closely related to the structural reduction, introduced in [14], which aims to find a short basis $\hat{B}$ of an overlattice $\mathcal{L}$ such that $\hat{L}/\mathcal{L}$ is isomorphic to some fixed abelian group $G$. However, the primary goal of [14] was to decrease the Gram–Schmidt norm of $\hat{B}$ in order to sample a pool of Gaussian overlattice vectors of norm $\Theta(\sqrt{n \log n \|B^*\|})$. These vectors would be too large for our purpose, since the bottom level of our decomposition algorithm needs a pool of vectors of length $\Theta(\sqrt{n \text{vol}(\mathcal{L})})$.

In the present paper, we prove that when the group $G$ is large enough, the unbalanced reduction of [14] can in fact efficiently construct a basis $C$ of $\mathcal{L}$ such that $[c_1/N^k, c_2, \ldots, c_n]$ is quasi-orthonormal. This naturally defines the tower of $k + 1$ overlattices $\mathcal{L}_i$, where $\mathcal{L}_i$ is generated by the corresponding basis $B^{(i)} = [c_1/N^i, c_2, \ldots, c_n]$ for $i = 0, \ldots, k$. Then the
Gaussian sampling algorithms on $\mathcal{L}_k$ can be replaced by Schnorr–Euchner enumeration, with or without pruning, using $B^{(k)}$, and thus, the norm of the overlattice vectors can be decreased to $r_n\sqrt{\operatorname{vol}(\mathcal{L}_k)}$. For completeness, we recall in the Appendix the pseudocode of the unbalanced reduction from [14]. Here, we prove that it allows us to produce a quasi-orthonormal basis. Compared to [14], we added the condition that $\sigma \leq \min \|b_i^*\|$ on the input parameters, and consequently, one of the test cases in the main loop of the algorithm in [14] never occurs.

Theorem 3.2 and its corollary below prove that running the unbalanced reduction with a smaller parameter $\sigma$ than is considered in [14] allows us to construct a quasi-orthonormal overlattice basis in polynomial time. Note that in [14], the goal was to find an overlattice basis whose Gram–Schmidt length was smaller than $\sigma$ without any constraints on their Rankin factors. Here, the first vector of the overlattice basis may be larger than $\sigma$; we just ensure that the $n$th root of the volume is exactly $\sigma$ and that all Rankin factors are polynomially bounded. Thus, the main equation (3.7) of Theorem 3.2 cannot be directly deduced from [14], and we provide a full proof in the Appendix.

**Theorem 3.2 (Unbalanced reduction).** Let $\mathcal{L}(B)$ be an $n$-dimensional integer lattice with an LLL-reduced basis $B = [b_1, \ldots, b_n]$. Let $\sigma$ be a target length $\leq \min \|b_i^*\|$. Algorithm B.1 outputs in polynomial time a basis $C$ of $\mathcal{L}$ satisfying

$$\|c_i^*\| \leq \sigma \quad \text{for all } i \in [2, n],$$

$$\|c_1^*\| \leq \sigma n \cdot \operatorname{vol}(\mathcal{L})/\sigma^n,$$

$$\frac{\sigma^{n+1-i}}{\operatorname{vol}(C_{[1,n]})} \leq n + 1 - i \quad \text{for all } i \in [2, n].$$

Since $\sigma$ is by construction the $n$th root of the bottom lattice $\mathcal{L}_k$, we immediately deduce the following elementary corollary, which proves that Algorithm 3 computes a tower of overlattices suitable for the decomposition algorithm.

**Corollary 1.** Given as input an LLL-reduced basis $B$ of $\mathcal{L}$ such that $\max \|b_i^*\|/\min \|b_i^*\| \leq 2^{O(n)}$, Algorithm 3 outputs a sequence of bases $B^{(0)}, \ldots, B^{(k)}$ such that $B^{(0)}$ is a basis of $\mathcal{L}$, $B^{(k)}$ is quasi-orthogonal, and $\mathcal{L}(B^{(i)})/\mathcal{L}(B^{(i-1)}) \simeq \mathbb{Z}/N\mathbb{Z}$ for all $i \in [1, k]$.

**Proof.** The condition $\max \|b_i^*\|/\min \|b_i^*\| \leq 2^{O(n)}$ ensures that the number of levels computed in Step 2 is linear in $n$. Thus, $\sigma$, computed in Step 3 is less than or equal to $\min \|b_i^*\|$. From the definition of $B^{(k)}$, we deduce that $\operatorname{vol}(\mathcal{L}(B^{(k)})) = \operatorname{vol}(\mathcal{L})/N^k = \sigma^n$, and for all $i \geq 2$, $B^{(i)}_{[i,n]} = B_{[i,n]}$. Thus, equation (3.7) proves that the Rankin factor $\gamma_{n,i}(B^{(k)}) = \sigma^{n-i}/\operatorname{vol}(B^{(k)}_{[i+1,n]})$ is less than or equal to $n - i$ for all $i \in [1, n - 1]$.

The proofs of Theorem 3.2 and Algorithm B.1 are given in Appendix B.

### 3.3. Cost for initial enumeration at level $k$ and pruning

The cost of a full enumeration of any bounded coset $(z + \mathcal{L}_k) \cap \operatorname{Ball}_n(r_n\beta\sigma)$ at level $k$ is:

$$T_{SE} = \sum_{i=1}^{n} \frac{\operatorname{vol}(\operatorname{Ball}(r_n\beta\sigma))}{\operatorname{vol}(B^{(k)}_{[n+1-i,n]})} \leq n \sum_{i=1}^{n} V_i \cdot (r_n\beta)^i = \tilde{O}(2^{0.398n})$$

where for $n \to \infty$ the maximal term in the sum, $\sim(\sqrt{n}/\sqrt{i})^i$, appears for $i = n\beta^2/e$. It is of size $\tilde{O}(2^{0.398n})$ because $\sqrt{e}^3/e \approx 2^{0.398}$. Experiments show that the above estimate is close to

---

1Equations (3.5) and (3.6) are in common with [14] and correspond more or less to the definition of the unbalanced reduction algorithm.
what we observe in practice, as we present in § 4. The number of steps in the full enumeration is an exponential factor less than $2^{0.03n}$ larger than the complexity of the merge. In practical dimensions up to 100, the actual running time of the full enumeration is already smaller than the time for the merge by collision in the consecutive steps, as elementary operations in the enumeration are faster than memory access and vector additions in the merge. However, more work must be done in large dimensions. For instance, a light pruning [12, 15] can be used to divide the running time of the initial enumeration by a small exponential factor of $2^{0.03n}$, but it will only recover a subset $S_k \subseteq C_k$. For instance, by use of the linear bounding function of [15], it will recover a fraction $1/n$ of $C_k$, and since depth $n+1-i$ in the pruned enumeration only explores a subset of the $i$-dimensional ball of radius $\sqrt{i/n}r_n\beta\sigma$ instead of $r_n\beta\sigma$, the running time $T_{\text{SE}}$ provably decreases below

$$T_{\text{lin,prun.}} \leq \sum_{i=1}^{n} \frac{\text{vol}(\text{Ball}_i(\sqrt{(i/n)r_n\beta\sigma}))}{\text{vol}(B_{i/n-1-i,n})} \leq n \sum_{i=1}^{n} V_i(r_n\sqrt{\frac{i}{n}}\beta)^i \approx n^2 \beta^n = \tilde{O}(2^{0.292n}). \quad (3.9)$$

Of course, there are lots of pruning tradeoffs between equations (3.8) and (3.9). This leads to a natural question on the stability of the algorithm: if the input of the merge at level $i$ is an incomplete subset $S_{i+1}$ containing only a constant or polynomially small fraction $\nu$ of all elements of $C_{i+1}$, is the merge algorithm still able to retrieve the whole set $C_i$? Intuitively, under some reasonable independence heuristics, $\beta$ should then be increased so that the volume of each ball intersection grows by a factor $1/\nu^2$. Thus condition (3.3) becomes $\beta^n \sqrt{1 - (\alpha/2)^{3H}} > \sqrt{nG_H/0.692\nu^2}$. On the other hand, $G_H$ can now be decreased from some large enough constant down to almost 1, since the Heuristic 2.1 only needs to be valid for a fraction at least $\nu$ of all intersections of balls, in order to get a fraction at least $\nu$ of $C_i$ in the output. Working with incomplete cosets also raises additional questions: how likely are short elements to be present in the incomplete output coset, and can this probability be increased with randomization and standard repetition arguments?

In the next section, we address these questions in our experimental results which implicitly uses $G_H = 1$ for efficiency reasons.

### 4. Experimental validation

In this section we present our experimental results of a C++ implementation of our Algorithm 1, presented in §3. We make use of the newNTL [16] and fplll [7] libraries as well as the Open MP [32] and GMP [11] library. We tested the algorithm on random lattices of dimensions up to $n = 90$ as input.

#### 4.1. Overview

Tests in smaller and larger dimensions confirm the choice of parameters $\alpha$ and $\beta$ that we computed for the asymptotic case. We are hence able to enumerate vectors of a target coset $C_0 = (t_0 + L_0) \cap \text{Ball}(R_0)$ and in this way we solve SVP as well as CVP. Indeed, unlike classical sieving algorithm, short elements, that is, either a short vector or a close vector, have a higher probability of being found than larger elements. Thus, even though we might miss some elements of the target coset, we almost always solve the respective SVP or CVP. For instance, the algorithm finds the same shortest vectors as solutions for the SVP challenges published in [35]. The memory requirement and running time in the course of execution closely match our estimates and the intermediate helper lattices $L_i$ behave as predicted.

Besides the search for one smallest/closest vector, each run of the algorithm, with appropriate parameters, finds a non-negligible fraction of the whole bounded coset $C_0$. Repeating the search for vectors in $C_0$ several times on a randomized LLL-reduced basis will discover the complete...
bounded coset. Our experiments reflect this behavior where we can use the Gaussian heuristic or Schnorr–Euchner enumeration to verify the proportion of recovered elements of $C_0$.

All these tasks can be performed by a single machine or independently by a cluster as a distributed computation.

4.2. Recovering $C_0$ in practice for smaller dimensions

For design reasons we have described an algorithm that produces the same number of elements per list in each iteration in order to find all of $C_0$. All lists contain $\#C_0 = \#((t_0 + L_0) \cap \text{Ball}_n(R_0)) \approx (1 + \varepsilon_n)^n \beta^n$ elements on average, where $\varepsilon_n$ can be neglected for very large dimensions (see also (3.3)). For accessible dimensions, we need to increase the radii of the balls slightly, by a small factor $1 + \varepsilon_n$, to compensate for small variations from the heuristic estimate. We here present results for different values $\varepsilon_n \leq 0.08$ and dimension $n \in \{40, 45, 50, 55, 60\}$. The larger the dimension, the better Heuristic 2.1 holds, which means that $\varepsilon_n$ can be chosen smaller (see (3.4)). Figure 4 shows the relation between varying $\varepsilon_n$ and the fraction of vectors of $C_0$ found for dimension $n \in \{40, 45, 50, 55, 60\}$. The optimal choice for $\varepsilon_n$ depends on $n$ and the fraction of $C_0$ we wish to enumerate.

4.3. Probability of success for randomized repetitions, example: small dimension

The success ratio of recovering all of $C_0$ rises with increasing $n$. We here present the case of smaller dimensions $n = \{50, 55\}$ to show how it evolves.

Suppose that we want to enumerate 100% of a coset $C_0$ in dimension 50. According to Figure 4, we need to choose $\varepsilon_n$ at least 0.07, which results in lists of size $(1 + \varepsilon_n)^n \beta^n \approx 29.4 \beta^n$ and a running time $(1 + \varepsilon_n)^{100} \beta^50 \approx 867.7$ on average. An alternative, which is less memory-consuming, is to choose a smaller $\varepsilon_n$, and to run the algorithm several times on randomized input bases. For instance, if one chooses $\varepsilon = 0.0535$, one should expect to recover $p = 6\%$ of $C_0$ per iteration on average. Then, assuming that the recovered vectors are uniformly and independently distributed in $C_0$, we expect to find a fraction of $1 - (1 - p)^r$ after $r$ repetitions.

To confirm this independence assumption, we tested repeated execution for SVP instances with parameters $n = 50$, $(1 + \varepsilon)\beta = 1.0535 \sqrt{3}/2$, $\alpha = \sqrt{4}/3$. Figure 5 shows the average number of distinct vectors of $C_0$ recovered as a function of the number of repetition $r$ (and the observed standard deviation) in comparison to the expected number of elements $C_0 \cdot (1 - (1 - 0.06)^r)$. The experiments match closely the estimate.


For a random lattice of dimension $n = 50$ and $\varepsilon = 0.0535$, the size of the coset $C_0$ is roughly 342 000. In our experiments, we found 164 662 vectors (48%) after 10 repetitions in which we randomized the basis. After 20 trials we found 239 231 elements which corresponds to 70%, and after 70 trials we found 337 016 elements (99% of $C_0$). We obtained the following results in dimension $n = 55$. After 10 trials with $\varepsilon = 0.0535$ we obtain 96.5% of the vectors of $C_0$, which is significantly higher in comparison to the 48% recovered after 10 trials in dimension 50.

4.4. Shorter or closer vectors are easier to find

During the merge operations, we can find a vector $v \in C_i$ if there exist vectors in the intersection between two balls of the same radius, centered at the end points of $v$. As the intersection is larger when $v$ is shorter (see Figure 6), we can deduce that with the practical variant, short vectors of a coset are easier to find than longer ones.

As we work with cosets, this means that vectors which are closer to the target (that is, short lattice vectors when the target is 0) should appear more often for different runs on a randomized input basis. We verified this observation experimentally by comparing the norm of a vector with the number of appearances during 100 repetitions in dimension 50, with $\varepsilon = 0.0535$, see Figure 7.
4.5. Parallelization

The algorithm itself is highly parallelizable for various types of hardware architectures. Of course, the dominant operations are \( n \)-dimensional vector additions and Euclidean norm computations, which can be optimized on any hardware containing vector instructions. Additionally, unlike sieving techniques, each iteration of the outer loop of the merge algorithm (Algorithm 2, line 3) can be run simultaneously, as every vector is treated independently of the output. Furthermore, one may divide the pool of vectors into \( p \leq \alpha^n/2 \) groups of buckets at each level, as soon as any two opposite buckets belong to the same group. Thus, the merge operation can operate on a group independently of all other groups. This allows us to efficiently run the algorithm when the available RAM is too small to store lists of size \( (1 + \varepsilon)^n \beta^n \). It also allows us to distribute the merge step on a cluster. For instance, in dimension \( n = 90 \) using \( \varepsilon = 0.0416 \), storing the full lists would require 3 TB of RAM. We divided the lists into 25 groups of 120 GB each, which we treated one at a time in RAM while the others were kept on hard drive. This did not produce any noticeable slowdown. Finally, the number of elements in each bucket can be estimated precisely in advance using Heuristic 2.1, and each group performs exactly the same vector operations (floating point addition, Euclidean norm computation) at the same time. This makes the algorithm suitable for SIMD implementation, not only multi-threading.

4.6. Experiments in low- and middle-sized dimensions

Our experiments in dimension 40–90 on challenges in [35] show that we find the same short vectors as those previously reported in the hall of fame. To solve SVP or CVP by use of the decomposition technique, it is in fact not necessary to enumerate the complete bounded coset \( C_0 \) and to ensure that the lists are always of size \( (1 + \varepsilon_n)^n \beta^n \) as we describe in the following paragraphs.

We give more details for medium dimensions \( n = 70 \) and \( n = 80 \) with \( \alpha = \sqrt{4/3} \) and \( \beta = \sqrt{3/2} \) in the following. The algorithm ran on a machine with an Opteron 6176 processor, containing 48 cores at 2.3 GHz, and having 256 GB of RAM. Table 2 presents the observed size of the lists \( S_i \subseteq C_i \) for each level in dimension 70 and 80.

In dimension 80, we chose aborted-BKZ-30 [17] for preprocessing. The algorithm has eight levels and we chose \( \varepsilon = 0.044 \) to obtain 97\% of \( C_0 \) after a single run. The initial enumeration on one core took a very short time of 6.5 CPU hours (so less than 10 min with our multi-thread
implementation of the enumeration) while each of the eight levels of the merge took between 20 and 36 CPU hours (so less than 45 min per level in our parallel implementation).

The number of elements in lower levels lies below the heuristic estimate and we keep loosing elements during the merge for the deepest levels. For example, in dimension 80 we start with 73% of $C_8$ and recover only 43% of $C_7$ after one step. Towards higher levels, we slowly begin to recover more and more elements. In dimension 80, the size of the lists starts to increase from level 5 on as $S_5, S_4$ and $S_3$ cover 41%, 47% and 56% of the vectors, respectively. This continues until the final step where we find 97% of the elements of $C_0$.

4.7. Pruning of the merge step in practice, larger dimension $n = 75$ and $n = 90$

In § 3.1, we obtain conditions on the parameters as we request the intersection $I$ of two balls to be non-empty, which means that $\frac{\text{vol}(I)}{\text{vol}(L)} \geq K$ for some number $K > 1$ under Heuristic 2.1. This condition suggests that at each level, each coset element in an output list $S_{i-1} \subseteq C_{i-1}$ of a merge is obtained on average about $K$ times. If the input list $S_i$ is shorter than expected, one will indeed recover fewer than $K$ copies of each element, but we may still have one representative of each element of $C_{i-1}$. Our experiments confirm this fact (see Tables 2 and 3). To solve SVP or CVP, one may shorten the time and memory necessary to find a solution vector by interrupting each level whenever the output list contains a sufficiently large fraction of the elements of the bounded cosets. For example, we ran our algorithm on the 75-dimensional basis of the SVP challenge [35] with seed 38. We chose $\varepsilon = 0.044$ and interrupted the merge if the size of the intermediate set $S_i$ reached 50% or 35% of $\#C_i$ for $i \in [1, k - 1]$. Table 3 presents the intermediate list sizes. In the end, we recovered 69% and 6.4% of $\#C_0$, respectively, and the shortest vector was found in both cases. The running time for the merge in the intermediate levels decreases compared to no pruning by a factor 0.49 and 0.29, respectively, as one would expect for lists that are smaller by at least a factor 0.5 and 0.35, respectively.

In dimension 90, we ran our algorithm on the 90-dimensional SVP challenge with seed 11, using $\varepsilon = 0.0416$. We chose to keep at most 33% of $C_i$ for $i \in [1, k - 1]$. Despite this harsh cut, the size of the intermediate lists remained stable after the first merge. And interestingly, after only 65 h on 32 threads, we recovered 61% of $\#C_0$ in the end, including the published shortest vector. Note that as we interrupt the merge, we in fact do not read all elements of the starting list $S_k$. One might hence simply not apply a full enumeration in practice but stop the Schnorr–Euchner enumeration once enough elements are enumerated.

4.8. Notes on the Gaussian heuristic for intermediate levels

Our quasi-orthogonal lattices at the bottom level behave randomly and follow the Gaussian heuristic. The most basic method to fill the bottom list $S_k$ is to run Schnorr–Euchner enumeration (see § 2) where the expected number of nodes in the enumeration tree is given by (3.8) based on Heuristic 2.1. Previous research has established that this estimate is accurate.
for random BKZ-reduced bases of random lattices in high dimension. Here, since we work with quasi-orthogonal bases, which are very specific, we redo the experiments, and confirm the findings also for quasi-orthogonal bases. Already for small dimensions \( n = 40, 50, 55 \), experiments show that the actual number of nodes in a Schnorr–Euchner enumeration is very close to the expected value. Figure 8 shows that the experiment and heuristic estimate for dimension 55, for example, are almost indistinguishable.

We also make use of Heuristic 2.1 when we estimate the number of coset vectors in the intersection of two balls. As the lower lattices in the tower are not ‘random’ enough, they have close to quasi-orthonormal bases, we observe smaller lists in the lower levels and thus a deviation from the heuristic. Beside the geometry of lattices, the deviation depends on the center of the balls or the center of the intersection. Randomly centered cosets of quasi-orthonormal lattices contain experimentally an average number of points a constant factor below \((1 + \varepsilon_n)^n \beta^n\). Zero-centered cosets contain more points, and should be avoided. The randomization of the initial target used in Algorithm 1 ensures that the centers are random modulo \( L_k \), even in an SVP setting. The number of vectors hence stays below but close to the estimate \((1 + \varepsilon_n)^n \beta^n\) after the first collision steps. The following steps can only improve the situation. The lattices in higher levels are more and more random and we observe that the algorithm recovers the expected number of vectors. This is a sign that our algorithm is stable even when the input pools \( S_i \) are incomplete.

Finally, experiments support the claim that the number of elements per bucket during the merge by collision corresponds to the average value \((\beta/\alpha)^n\). For example, in dimension \( n = 80\),

<table>
<thead>
<tr>
<th>Level = ( i )</th>
<th>9</th>
<th>8</th>
<th>7</th>
<th>6</th>
<th>5</th>
<th>4</th>
<th>3</th>
<th>2</th>
<th>1</th>
<th>0</th>
<th>SVP</th>
</tr>
</thead>
<tbody>
<tr>
<td>( n = 75, )</td>
<td></td>
<td></td>
<td>50</td>
<td>50</td>
<td>47</td>
<td>46</td>
<td>46</td>
<td>48</td>
<td>50</td>
<td>69</td>
<td></td>
</tr>
<tr>
<td>( \varepsilon = 0.044 )</td>
<td>Heuris.</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Solved</td>
<td></td>
</tr>
<tr>
<td>( n = 75, )</td>
<td></td>
<td></td>
<td>50</td>
<td>35</td>
<td>30</td>
<td>25</td>
<td>20</td>
<td>15</td>
<td>8</td>
<td>6.4</td>
<td></td>
</tr>
<tr>
<td>( \varepsilon = 0.044 )</td>
<td>Heuris.</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Solved</td>
<td></td>
</tr>
<tr>
<td>( n = 90, )</td>
<td></td>
<td></td>
<td>70</td>
<td>40</td>
<td>40</td>
<td>40</td>
<td>40</td>
<td>40</td>
<td>40</td>
<td>70</td>
<td></td>
</tr>
<tr>
<td>( \varepsilon = 0.0416 )</td>
<td>Heuris.</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Solved</td>
<td></td>
</tr>
<tr>
<td>( n = 90, )</td>
<td></td>
<td></td>
<td>70</td>
<td>33</td>
<td>33</td>
<td>33</td>
<td>33</td>
<td>33</td>
<td>33</td>
<td>61</td>
<td></td>
</tr>
<tr>
<td>( \varepsilon = 0.0416 )</td>
<td>Heuris.</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Cut</td>
<td>Solved</td>
<td></td>
</tr>
</tbody>
</table>
for parameters $\alpha = \sqrt{4/3}$, $\beta = \sqrt{3/2}$, $\varepsilon = 0.044$, we observe that the largest bucket contains only 10% more elements than the average value, and that 60% of the buckets are within $\pm 2\%$ of the average value.

4.9. Comparison to experimental results of a parallel Gauss sieve algorithm

From a very general point of view, our algorithm can be viewed as a sieving algorithm. The algorithm is decomposed into a polynomial number of levels, each one corresponds to a certain upper bound $R_i$ on the norm. At each level, we use an exponential pool of lattice vectors, perform linear combinations, and select the shortest of them for the next level.

We now list the specificities of our algorithm compared to previous sieving algorithms:

- We start from short vectors (in overlattices), and at each level the norm $R_i$ geometrically increases by a factor $\alpha$.
- At each level, we maintain an (almost) complete set of all coset vectors of norm less than or equal to $R_i$. For this reason, our algorithm has the stability property that short coset vectors are more likely to be found. Classical sieving techniques satisfy the opposite: short vectors have a negligible probability of appearing spontaneously. Our algorithm is compatible with pruning, and it can solve the Exact CVP. Reducing the list sizes for classical sieving leads in general to catastrophic results.
- Our algorithm is highly parallelizable as it allows us to use up to $\alpha^n$ independent threads per merge operation as explained in §4.5. However, the accessible dimension is naturally limited by the exponential memory requirement of order $\beta^n$. There exists parallel versions of the Gauss sieve [19, 29], which leads to faster practical running time in dimensions 70 to 96, however the efficiency of the parallelization decreases fast when the number of threads increases because of the list size and the communication cost [19].
- Our algorithm is essentially a CVP solver and is not specialized for SVP: if classical sieving algorithms were to be turned into CVP solvers, then it would obviously be impossible to regroup each vector with its opposite, and the lists of vectors would be twice as large. Furthermore, classical sieving techniques rely on the fact that vectors which cannot be reduced by others, necessarily become poles to reduce others. By replacing subtractions with additions in order to preserve the target, these two options, reducing a vector $v$ by others and considering $-v$ as a pole, cease to be mutually exclusive, and both would have to be tested. Thus, turning classical sieving algorithms into CVP solvers would likely increase their running time by a factor of 4 and their memory requirement by a factor of 2, with no guarantee that they would actually find the solution.

We give some concrete timings. To solve instances in dimension 80 and 90, our algorithm takes more time than the currently fastest implementation of the Gauss sieve algorithm [19]. Ishiguro et al. report in [19] solving the SVP challenge in dimension 80 in 29 sequential hours and an instance of dimension 96 in 6400 sequential hours. Our algorithm, however, needs 65 sequential hours in dimension 80 and 2080 hours in dimension 90. It is slower than the Gauss sieve used as an SVP solver, yet the slowdown factor remains smaller than 4, which would be expected (as a minimum) to turn it into a CVP solver.

5. Conclusion

We have presented an alternative approach to solve the hard lattice problems SVP and CVP for random lattices. It makes use of a new technique that is different from those used so far in enumeration or sieving algorithms, and works by moving short vectors along a tower of nested lattices. Our experiments show that the method works well in practice. An open question in the case of ideal lattices is to find a structural reduction that preserves the cyclic structure or provides an other structure for which speed-ups can be applied.
Appendix A. Intersection of hyperballs

The volume of the intersection, $\text{vol}_I(d)$, of two $n$-dimensional hyperballs of radius 1 at distance $d \in [0.817; 2]$ can be approximated for large $n$ by the volume of the $n$-dimensional ball of radius $D = \sqrt{1 - (d/2)^2}$ (see Lemma A.1 below). If we consider the intersection of two balls of radius $R$, the volume gets multiplied by a factor of $R^n$, as stated in Corollary A.2.

**Lemma A.1.** The volume of the intersection of two $n$-dimensional hyperballs of radius 1 at distance $d \in [0.817; 2]$ is

$$\frac{2V_{n-1}}{(n+1)V_n} \arccos\left(\frac{d}{2}\right) \leq \frac{\text{vol}_I(d)}{\text{vol}(\text{Ball}_n(D))} \leq \frac{2V_{n-1}}{(n/2 + 1)V_n} \arccos\left(\frac{d}{2}\right)$$

where $D = \sqrt{1 - (d/2)^2}$.

**Proof.** The intersection of two balls of radius 1 whose centers are at distance $d \in [0, 2]$ of each other can be expressed as

$$\text{vol}_I(d) = 2 \cdot \int_{d/2}^{1} V_{n-1}(\sqrt{1 - x^2})^{n-1} dx = 2V_{n-1} \int_{0}^{\arccos(d/2)} \sin^n(\theta) d\theta$$

where $V_{n-1}$ equals the volume of the $(n-1)$-dimensional ball of radius 1. For $d \in [0.817; 2]$ one can bound the sine term in the integral:

$$\frac{D}{\arccos(d/2)} \theta \leq \sin(\theta) \leq \frac{D}{\sqrt{\arccos(d/2)}} \sqrt{\theta}.$$

Therefore, we obtain bounds for the volume of the intersection,

$$\text{vol}_I(d) \leq \frac{2V_{n-1}}{n/2 + 1} \arccos\left(\frac{d}{2}\right) D^n$$

and

$$\text{vol}_I(d) \geq \frac{2V_{n-1}}{n + 1} \arccos\left(\frac{d}{2}\right) D^n,$$

which proves the lemma.

We can use the lower bound of Lemma A.1 and obtain a numerical lower bound on the volume of the intersection of balls of radius $R$ at distance at most $\sqrt{4/3}R$ used in our algorithm:

**Corollary A.2.** For all dimensions $n \geq 10$, the volume of the intersection of two $n$-dimensional hyperballs of radius $R$ at distance $dR$, where $d \leq \sqrt{4/3}$, is lower-bounded by

$$R^n \text{vol}_I(d) \geq \frac{0.692}{\sqrt{n}} \cdot R^n \text{vol}(\text{Ball}_n(\sqrt{1 - (d/2)^2})).$$

Appendix B. Proof of Theorem 3.2 and Algorithm B.1

We use the suffixes ‘old’ and ‘new’ to denote the values of the variables at the beginning and end of the ‘for’ loop of Algorithm B.1, respectively. Furthermore, we call $x_i$ the value $\|b_i^{\text{new}}\|$ during iteration $i$. Note that $x_i$ is also $\|b_i^{\text{old}}\|$ during the next iteration (of index $i - 1$ since $i$ goes backwards).

For $i \in [1, n]$, let $a_i = \|b_i^{\text{old}}\|/\sigma$. Note that $a_i$ is always 1 or greater. We show by induction over $i$ that the following invariant holds at the end of each iteration of Algorithm B.1:
An LLL-reduced basis

Input: A basis $C$ of $\mathcal{L}$ satisfying $\|b_1\| \leq \sigma \text{vol}(\mathcal{L})/\sigma^n$, and, for all $i \in [2, n]$, $\|c_i^*\| \leq \sigma$ and $\frac{\sigma^{n+1-i}}{\text{vol}(C_{[i,n]})} \leq n + 1 - i$.

Output: A basis $C$ of $\mathcal{L}$ satisfying $\|c_1\| \leq \sigma \text{vol}(\mathcal{L})/\sigma^n$, and, for all $i \in [2, n]$, $\|c_i^*\| \leq \sigma$ and $\frac{\sigma^{n+1-i}}{\text{vol}(C_{[i,n]})} \leq n + 1 - i$.

Algorithm B.1 Unbalanced reduction from [14], specialized for $\sigma \leq \min \|b_i^*\|$.

1: $C \leftarrow B$
2: Compute the Gram–Schmidt matrices $\mu$ and $C^*$
3: Let $k$ be the largest index such that $\|c_k^*\| > \sigma$
4: for $i = k - 1, \ldots, 1$ do
5: \hspace{1em} $\gamma \leftarrow -\mu_{i+1,i} + \frac{\|c_{i+1}^*\|}{\|c_i^*\|} \sqrt{(\frac{\|c_i^*\|}{\sigma})^2 - 1}$
6: \hspace{1em} $(c_i, c_{i+1}) \leftarrow (c_{i+1} + \gamma \cdot c_i, c_i)$
7: Update the Gram–Schmidt matrices $\mu$ and $C^*$
8: end for
9: return $C$

At the first iteration ($i = k - 1$), it is clear that $x_k = \|b_k^{\text{old}}\| = \sigma a_k$. At the beginning of iteration $i$, we always have $\|b_i^{\text{old}}\| > \sigma$, and by induction, $\|b_i^{\text{old}}\| > \sigma$. We transform the block so that the norm of the first vector satisfies

$$R \leq \|b_i^{\text{new}}\| \leq R + \|b_i^{\text{old}}\|,$$

where $R = \|b_i^{\text{old}}\|/\|b_i^{\text{old}}\|/\sigma$.

This condition can always be fulfilled with a primitive vector of the form $b_i^{\text{new}} = b_i^{\text{old}} + \gamma b_i^{\text{old}}$ for some $\gamma \in \mathbb{Z}$. Since the volume is invariant, the new $\|b_i^{\text{new}}\|$ is upper-bounded by $\sigma$.

By developing (B.1), we derive a bound on $x_1$,

$$x_1 \leq \sigma \sum_{i=1}^k a_1 \ldots a_i \leq \sigma \sigma \prod_{i=1}^k a_i \leq n \sigma \text{vol}(\mathcal{L})/\sigma^n,$$

which proves (3.6). Similarly, one obtains that $x_i \leq (n + 1 - i) \sigma \text{vol}(B_{[i,n]})/\sigma^{n+1-i}$, which is equivalent to (3.7). Note that the transformation matrix of the unbalanced reduction algorithm is

$$
\begin{bmatrix}
\gamma_1 & \ldots & \gamma_{k-1} & 1 & 0 & \ldots & 0 \\
1 & 0 & \ldots & 0 & \vdots & \vdots & \\
0 & \ddots & \ddots & \vdots & \vdots & \vdots & \\
0 & 0 & 1 & 0 & 0 & \ldots & 0 \\
0 & \ldots & \ldots & 0 & 1 & 0 & 0 \\
\vdots & \vdots & \ddots & 0 & \ddots & \vdots & \\
0 & \ldots & \ldots & 0 & 0 & 0 & 1
\end{bmatrix},
$$

where $\gamma_i$ is $[-\mu_{i+1,i} + x_{i+1}/\sqrt{1 - 1/a_i^2}]$. Since each $x_{i+1}$ is bounded by
all coefficients have a size polynomial in the input basis. This proves that Algorithm B.1 has polynomial running time. \hfill \qed

References

7. X. Cadé, D. Pujol and D. Stehlé, fplll 4.0.4, May 2013.

Anja Becker
EPFL, École Polytechnique Fédérale de Lausanne
Laboratory for Cryptologic Algorithms (LACAL)
Switzerland
anja.becker@epfl.ch

Nicolas Gama
UVSQ/PRISM
Université de Versailles
France
nicolas.gama@prism.uvsq.fr

Antoine Joux
CryptoExperts
INRIA/Ouragan
Chaire de Cryptologie de la Fondation de l’UPMC
Sorbonne Universités
UPMC Univ Paris 06
CNRS UMR 7606
LIP 6
France
Antoine.Joux@m4x.org