# Parallel Enumeration of Shortest Lattice Vectors 

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#### Abstract

Lattice basis reduction is the problem of finding short vectors in lattices. The security of lattice based cryptosystems is based on the hardness of lattice reduction. Furthermore, lattice reduction is used to attack well-known cryptosystems like RSA. One of the algorithms used in lattice reduction is the enumeration algorithm (ENUM), that provably finds a shortest vector of a lattice. We present a parallel version of the lattice enumeration algorithm. Using multi-core CPU systems with up to 16 cores, our implementation gains a speed-up of up to factor 14. Compared to the currently best public implementation, our parallel algorithm saves more than $90 \%$ of runtime.


Keywords: lattice reduction, shortest vector problem, cryptography, parallelization, enumeration

## 1 Introduction

A lattice $\mathcal{L}$ is a discrete subgroup of the space $\mathbb{R}^{d}$. Lattices are represented by linearly independent basis vectors $\mathbf{b}_{1}, \ldots, \mathbf{b}_{n} \in \mathbb{R}^{d}$, where $n$ is called the dimension of the lattice. Lattices have been known in number theory since the eighteenth century. They already appear when Lagrange, Gauss, and Hermite study quadratic forms. Nowadays, lattices and hard problems in lattices are widely used in cryptography as the basis of promising cryptosystems.

One of the main problems in lattices is the shortest vector problem (SVP), that searches for a vector of shortest length in the lattice. The shortest vector problem is known to be NP-hard under randomized reductions. It is also considered to be intractable even in the presence of quantum computers. Therefore, most lattice based cryptographic primitives, e.g. one-way functions, hash functions, encryption, and digital signatures, leverage the complexity of the SVP problem. In the field of cryptanalysis, lattice reduction is used to attack the NTRU and GGH cryptosystems. Further, there are attacks on RSA and low density knapsack cryptosystems.

Lattice reduction still has applications in other fields of mathematics and number theory. It is used for factoring composite numbers and computing discrete logarithms using diophantine approximations [Sch91]. In the field of dis-
crete optimization, lattice reduction is used to solve linear integer programs [Len83].

The fastest algorithm known to solve SVP is the enumeration algorithm of Kannan [Kan83] and the algorithm of Fincke and Pohst [FP83]. The variant used mostly in practice was presented by Schnorr and Euchner in 1991 [SE94]. Nevertheless, these algorithms solve SVP in exponential runtime. Therefore, enumeration is only applicable in low lattice dimensions $(n \leq 60)$. For higher dimensions it is only possible to find short vectors, but not a shortest vector. Mostly, these approximate solutions of the SVP are sufficient in practice. In 1982 the famous LLL algorithm was presented for factoring polynomials [LLL82]. This algorithm does not solve SVP rather it finds a vector with length exponential in the lattice dimension. However, LLL is the first algorithm having a polynomial asymptotic running time. LLL can be run in lattice dimension up to 1000.

In practice, the most promising algorithm for lattice reduction in high dimensions is the BKZ block algorithm by Schnorr and Euchner [SE94]. It mainly consists of two parts, namely enumeration in blocks of small dimension and LLL in high dimension. BKZ finds shorter vectors than LLL, at the expense of a higher runtime.

Considering parallelization, there are various works dealing with LLL, e.g., [Vil92,BW09]. For the more time-consuming part of BKZ, namely the enumeration step, assuming the use of high block sizes, there is no work published about parallelizing.

The enumeration in lattices can be visualized as a depth first search in a weighted search tree, with different subtrees being independent from each other. Therefore, it is possible to enumerate different subtrees in parallel threads without any communication between threads. This makes enumeration a good candidate for platforms that act in a MIMD (multiple instruction - multiple data) manner. Therefore, we have chosen multi-core CPUs for the implementation of our parallel enumeration algorithm.

Our Contribution. In this paper we parallelize the enumeration algorithm by Schnorr and Euchner [SE94]. We implement the parallel version of ENUM and test it on multi-core CPUs. More precisely, we use up to 16 CPU cores to speed up the lattice enumeration, in lattice dimensions of 40 and above. Considering the search tree, the main problem is to predict the subtrees that are examined during enumeration beforehand.

We gain speed-ups of up to factor 14 compared to our single core version. ${ }^{3}$ Compared to the fastest single-core ENUM implementation known our parallel version of ENUM saves more than $90 \%$ of runtime. We add some clever additional communication among threads, such that by using $s$ processor cores we even gain a speed-up of more than $s$ in some cases.

With this work we show that it is possible to parallelize the entire BKZ algorithm for the search for short lattice vectors. The strength of BKZ is used

[^0]to assess the practical hardness of lattice reduction, which helps finding suitable parameters for secure lattice based cryptosystems.

Organization. Section 2 explains the required basic facts on lattices and parallelization, Section 3 describes the ENUM algorithm by [SE94], Section 4 presents our new algorithm for parallel enumeration, and Section 5 gives our experimental results.

## 2 Preliminaries

Notation. Vectors and matrices are written in bold face, e.g. v and M. The expression $\lceil x\rfloor$ denotes the nearest integer to $x \in \mathbb{R}$, i.e., $\lceil x\rfloor=\lceil x-0.5\rceil$. $\|\mathbf{v}\|$ denotes the Euclidean norm, other norms are indexed with a subscript, like $\|\mathbf{v}\|_{\infty}$. Throughout the paper $n$ denotes the lattice dimension.

Lattices. A lattice is a discrete additive subgroup of $\mathbb{R}^{d}$. It can be represented as the linear integer span of $n \leq d$ linear independent vectors $\mathbf{b}_{1}, \ldots, \mathbf{b}_{n} \in \mathbb{R}^{d}$, which are arranged in a column matrix $\mathbf{B}=\left[\mathbf{b}_{1}, \ldots, \mathbf{b}_{n}\right] \in \mathbb{R}^{d \times n}$. The lattice $\mathcal{L}(\mathbf{B})$ is the set of all linear integer combinations of the basis vectors $\mathbf{b}_{i}$, namely $\mathcal{L}(\mathbf{B})=\left\{\sum_{i=1}^{n} x_{i} \mathbf{b}_{i}: x_{i} \in \mathbb{Z}\right\}$. The dimension of the lattice equals the number of linearly independent basis vectors $n$. If $n=d$, the lattice is called full-dimensional. For $n \geq 2$ there are infinitely many bases of a lattice. One basis can be transformed into another using a unimodular transformation matrix. The first successive minimum $\lambda_{1}(\mathcal{L}(\mathbf{B}))$ is the length of a shortest non-zero vector of a lattice. There exist multiple shortest vectors of a lattice, a shortest vector is not unique. Define the Gram-Schmidt-orthogonalization $\mathbf{B}^{*}=\left[\mathbf{b}_{1}^{*}, \ldots, \mathbf{b}_{n}^{*}\right]$ of $\mathbf{B}$. It is computed via $\mathbf{b}_{i}^{*}=\mathbf{b}_{i}-\sum_{j=1}^{i-1} \mu_{i, j} \mathbf{b}_{j}^{*}$ for $i=1, \ldots, n$, where $\mu_{i, j}=\mathbf{b}_{i}^{T} \mathbf{b}_{j}^{*} /\left\|\mathbf{b}_{j}^{*}\right\|^{2}$ for all $1 \leq j \leq i \leq n$. We have $\mathbf{B}=\mathbf{B}^{*} \mu^{T}$, where $\mathbf{B}^{*}$ is orthogonal and $\mu^{T}$ is an upper triangular matrix. Note that $\mathbf{B}^{*}$ is not necessarily a lattice basis.

Lattice Problems and Algorithms. The most famous problem in lattices is the shortest vector problem (SVP). The SVP asks to find a shortest vector in the lattice, namely a vector $\mathbf{v} \in \mathcal{L} \backslash\{0\}$ with $\|\mathbf{v}\|=\lambda_{1}(\mathcal{L})$. An approximation version of the SVP was solved by Lenstra, Lenstra, and Lovász [LLL82]. The LLL algorithm is still the basis of most algorithms used for basis reduction today. It runs in polynomial time in the lattice dimension and outputs a so-called $L L L$ reduced basis. This basis consists of nearly orthogonal vectors, and a short, first basis vector with approximation factor exponential in the lattice dimension. The BKZ algorithm by Schnorr and Euchner [SE94] reaches better approximation factors, but still has runtime exponential in the lattice dimension. Nevertheless, BKZ is the algorithm used mostly in practice today. As a subroutine, BKZ makes use of an exact SVP solver, such as ENUM. Theoretically, the best results for the approximation version of SVP are promised by the algorithm of [GN08a]. In practice, SVP can only be solved in low dimension $n$, say up to 60 , using
exhaustive search techniques or, a second approach, using sieving algorithms that work probabilistically. An overview of enumeration algorithms is presented in [PS08]. A randomized sieving approach for solving exact SVP was presented in [AKS01] and an improved variant in [MV10]. As the storage required by sieving algorithms is exponential in the lattice dimension, those algorithms are considered impractical. In this paper, we only deal with enumeration algorithms.

Exhaustive Search. In [PS08] Pujol and Stehlé examine the floating point behaviour of the ENUM algorithm. They state that double precision is suitable for lattice dimensions up to 90 , which is more than enumeration algorithms can handle in practice today. It is common practice to pre-reduce lattices before starting enumeration, as this reduces the radius of the search space. In BKZ, the basis is always reduced with the LLL algorithm when starting enumeration.

Publicly available implementations of enumeration algorithms are the established implementation of Shoup's NTL library and the fpLLL library of Stehlé et.al. Experimental data on enumeration algorithms using NTL can be found in [GN08b,MV10], both using NTL's enumeration.

Parallelization. Before we present our parallel enumeration algorithm, we need to introduce definitions specifying the quality of the realized parallelization. Furthermore, we give an overview of parallel computing paradigms.

There exist many parallel environments to perform operations concurrently. These models differ in their area of application. The most important parallel paradigms are the following:

MIMD (Multiple Instruction - Multiple Data): MIMD systems work in an asynchronous way, i.e. each processor performs its own operations (instructions). Therefore, they require communication to exchange information. They also might share a common memory.
SIMD (Single Instruction - Multiple Data): SIMD systems handle more processors compared to MIMD. Due to the simplicity of these processors, each processor runs the same operations (instructions), but with probably different inputs. Compared to MIMD they require less communication overhead, but also perform slower for arithmetic operations.

A multi-core microprocessor deploys the MIMD paradigm. Nowadays, such computer systems are commonly available. They possess several cores, while each core acts as an independent processor unit. The operating system is responsible to deliver operations to the cores.

In the parallelization context there exist notions that measure the achieved quality of a parallel algorithm compared to the sequential version. In the sequel of this paper, we will need the following definitions:

Speed-up factor: time needed for serial computation divided by the time required for the parallel algorithm. Using $s$ processes, a speed-up factor of up to $s$ is expected.

Efficiency: speed-up factor divided by the number of used processors. An efficiency of 1.0 means that $s$ processors lead to a speed-up factor of $s$ which can be seen as a "perfect" parallelization. Normally the efficiency is smaller than 1.0 because of the communication overhead for inter-process communication.

A parallel algorithm may benefit from communication, in such a way that less operations need to be computed. As soon as the number of saved operations exceeds the communication overhead, an efficiency of more than 1.0 might be achieved.

## 3 Enumeration of the shortest lattice vector

In this chapter we give an overview of the ENUM algorithm first presented in [SE94]. In the first place, the algorithm was proposed as a subroutine in the BKZ algorithm, but ENUM can be used as a stand-alone instance to solve the exact SVP. An example instance of ENUM in dimension 3 is shown by the solid line of Figure 1. An algorithm listing is shown as Algorithm 1.

```
Algorithm 1: Basic Enumeration Algorithm
    Input: Gram-Schmidt coefficients \(\left(\mu_{i, j}\right)_{1 \leq j \leq i \leq n},\left\|\mathbf{b}_{1}^{*}\right\|^{2} \ldots\left\|\mathbf{b}_{n}^{*}\right\|^{2}\)
    Output: \(\mathbf{u}_{\text {min }}\) such that \(\left\|\sum_{i=1}^{n} u_{i} \mathbf{b}_{i}\right\|=\lambda_{1}(\mathcal{L}(\mathbf{B}))\)
    \(A \leftarrow\left\|\mathbf{b}_{1}^{*}\right\|^{2}, \mathbf{u}_{\text {min }} \leftarrow(1,0, \ldots, 0), \mathbf{1} \leftarrow(0, \ldots, 0), \mathbf{c} \leftarrow(0, \ldots, 0)\)
    \(t=1\)
    while \(t \leq n\) do
        \(l_{t} \leftarrow l_{t+1}+\left(u_{t}+c_{t}\right)^{2}\left\|\mathbf{b}_{t}^{*}\right\|^{2}\)
        if \(l_{t}<A\) then
            if \(t>1\) then
                \(t \leftarrow t-1 \quad \triangleright\) move one layer down in the tree
                \(c_{t} \leftarrow \sum_{i=t+1}^{n} u_{i} \mu_{i, t}, \quad u_{t} \leftarrow\left\lceil c_{t}\right\rfloor\)
            else
                \(A \leftarrow l_{t}, \quad \mathbf{u}_{\text {min }} \leftarrow \mathbf{u} \quad \triangleright\) set new minimum
        else
            \(t \leftarrow t+1 \quad \triangleright\) move one layer up in the tree
            choose next value for \(u_{t}\) using the zig-zag pattern
```

To find a shortest non-zero vector of a lattice $\mathcal{L}(\mathbf{B})$ with $\mathbf{B}=\left[\mathbf{b}_{1}, \ldots, \mathbf{b}_{n}\right]$, ENUM takes as input the Gram-Schmidt coefficients $\left(\mu_{i, j}\right)_{1 \leq j \leq i \leq n}$, the quadratic norm of the Gram-Schmidt orthogonalization $\left\|\mathbf{b}_{1}^{*}\right\|^{2}, \ldots,\left\|\mathbf{b}_{n}^{*}\right\|^{2}$ of $\mathbf{B}$, and an initial bound $A$. The search space is the set of all coefficient vectors $\mathbf{u} \in \mathbb{Z}^{n}$ that satisfy $\left\|\sum_{t=1}^{n} u_{t} \mathbf{b}_{t}\right\| \leq A$. Starting with an LLL-reduced basis, it is common to set $A=\left\|\mathbf{b}_{1}^{*}\right\|^{2}$ in the beginning. If the norm of the shortest vector is known beforehand, it is possible to start with a lower $A$, which limits the search space and reduces the runtime of the algorithm. If a vector $\mathbf{v}$ of length smaller than $A$ is found, $A$ can be reduced to the norm of $\mathbf{v}$, that means $A$ always denotes the size of the current shortest vector.

The goal of ENUM is to find a coefficient vector $\mathbf{u} \in \mathbb{Z}^{n}$ satisfying the equation

$$
\begin{equation*}
\left\|\sum_{t=1}^{n} u_{t} \mathbf{b}_{t}\right\|=\min _{x \in \mathbb{Z}^{n}}\left\|\sum_{t=1}^{n} x_{t} \mathbf{b}_{t}\right\| . \tag{1}
\end{equation*}
$$

Therefore, all coefficient combinations $\mathbf{u}$ that determine a vector of norm less than $A$ are enumerated. In Equation 1 we replace all $\mathbf{b}_{t}$ by their orthogonalization, i.e., $\mathbf{b}_{t}=\mathbf{b}_{t}^{*}+\sum_{j=1}^{t-1} \mu_{t, j} \mathbf{b}_{j}^{*}$ and get Equation (2):

$$
\left\|\sum_{t=1}^{n} u_{t} \mathbf{b}_{t}\right\|^{2}=\left\|\sum_{t=1}^{n}\left(u_{t} \cdot\left(\mathbf{b}_{t}^{*}+\sum_{j=1}^{t-1} \mu_{t, j} \mathbf{b}_{j}^{*}\right)\right)\right\|^{2}=\sum_{t=1}^{n}\left(u_{t}+\sum_{i=t+1}^{n} \mu_{i, t} u_{i}\right)^{2} \cdot\left\|\mathbf{b}_{t}^{*}\right\|^{2} .
$$

Let $\mathbf{c} \in \mathbb{R}^{d}$ with $c_{t}=\sum_{i=t+1}^{n} \mu_{i, t} u_{i}$ (line 8 ), which is predefined by all coefficients $u_{i}$ with $n \geq i>t$. The intermediate norm $l_{t}$ (line 4) is defined as $l_{t}=l_{t+1}+\left(u_{t}+c_{t}\right)^{2}\left\|\mathbf{b}_{t}^{*}\right\|^{2}$. This is the norm part of Equation 2 that is predefined by the values $u_{i}$ with $n \geq i \geq t$.

The algorithm enumerates the coefficients in reverse order, from $u_{n}$ to $u_{1}$. This can be considered as finding a minimum in a weighted search tree. The height of the tree is uniquely determined by the dimension $n$. The root of the tree denotes the coefficient $u_{n}$. The coefficient values $u_{t}$ for $1 \leq t \leq n$ determine the values of the vertices of depth $(n-t+1)$, leafs of the tree contain coefficients $u_{1}$. The inner nodes represent intermediate nodes, not complete coefficient vectors, i.e., a node on level $t$ determines a subtree $\left(\perp, \ldots, \perp, u_{t}, u_{t+1}, \ldots, u_{n}\right)$, where the first $t-1$ coefficients are not yet set. $l_{t}$ is the norm part predefined by this inner node on level $t$. We only enumerate parts of the tree with $l_{t}<A$. Therefore, the possible values for $u_{t}$ on the next lower level are in an interval around $c_{t}$ with $\left(u_{t}+c_{t}\right)^{2}<\left(A-l_{t+1}\right) /\left\|\mathbf{b}_{t}^{*}\right\|$, following the definition of $l_{t}$.

ENUM iterates over all possible values for $u_{t}$, as long as $l_{t} \leq A$, the current minimal value. If $l_{t}$ exceeds $A$, enumeration of the corresponding subtree can be cut off, the intermediate norm $l_{t}$ will only increase when stepping down in the tree, as $l_{t} \leq l_{t-1}$ always holds. The iteration over all possible coefficient values is (due to Schnorr and Euchner) performed in a zig-zag pattern. The values for $u_{t}$ will be sequenced like either $c_{t}, c_{t}+1, c_{t}-1, c_{t}+2, c_{t}-2, \ldots$ or $c_{t}, c_{t}-1, c_{t}+1, c_{t}-2, c_{t}+2, \ldots$

ENUM starts at the leaf $(1,0, \ldots, 0)$ and gives the first possible solution for a shortest vector in the given lattice. The algorithm performs its search by moving up (when a subtree can be cut off due to $l_{t} \geq A$ ) and down in the tree (lines 12 and 7). The norm of leaf nodes is compared to $A$. If $l_{1} \leq A$, it stores $A \leftarrow l_{1}$ and $\mathbf{u}_{\text {min }} \leftarrow \mathbf{u}$ (line 10), which define the current shortest vector and its size. When ENUM moves up to the root of the search tree it terminates and outputs the computed global minimum $A$ and the corresponding shortest vector $\mathbf{u}_{\text {min }}$.

## 4 Algorithm for parallel enumeration of the shortest lattice vector

In this section we describe our parallel algorithm for enumeration of the shortest lattice vector. The algorithm is a parallel version of the algorithm presented in [SE94]. First we give the main idea of parallel enumeration. Secondly, we present a high level description. Algorithms 2 and 3 depict our parallel ENUM. Thirdly, we explain some improvements that speed up the parallelization in practice.

### 4.1 Parallel Lattice Enumeration

The main idea for parallelization is the following. Different subtrees of the complete search tree are enumerated in parallel independently from each other representing them as threads (Sub-ENUM threads). Using $s$ processors, $s$ subtrees can be enumerated at the same time. All threads ready for enumeration are stored in a list L, and each CPU core that has finished enumerating a subtree picks the next subtree from the list. Each of the subtrees is an instance of SVP in smaller dimension; the initial state of the sub-enumeration can be represented by a tuple ( $\mathbf{u}, l, c, t$ ).

When the ENUM algorithm increases the level in the search tree, the center $\left(c_{t}\right)$ and the range $\left(\left(A-l_{t+1}\right) /\left\|\mathbf{b}_{t}^{*}\right\|\right)$ of possible values for the current index are calculated. Therefore, it is easy to open one thread for every value in this range.


Fig. 1. Comparison of serial (solid line) and parallel (dashed line) processing of the search tree.

Figure 1 shows a 3-dimensional example and compares the flow of the serial ENUM with our parallel version. Beginning at the starting node the procession order of the serial ENUM algorithm follows the directed solid edges to the root. In the parallel version dashed edges represent the preparation of new Sub-ENUM
threads which can be executed by a free processor unit. Crossed-out edges point out irrelevant subtrees. Threads terminate as soon as they reach either a node of another thread or the root node.

Extra Communication - Updating the Shortest Vector. Again, we denote the current minimum, the global minimum, as $A$. In our parallel version, it is the global minimum of all threads. As soon as a thread has found a new minimum, the Euclidean norm of this vector is written back to the shared memory, i.e. $A$ is updated. At a certain point every thread checks the global minimum whether another thread has updated $A$ and, if so, uses the updated one. The smaller $A$ is, the faster a thread terminates, because subtrees that exceed the current minimum can be cut off in the enumeration. The memory access for this update operation is minimal, only one integer value has to be written back or read from shared memory. This is the only type of communication among threads, all other computations can be performed independently without communication overhead.

### 4.2 The Algorithm for Parallel Enumeration

Algorithm 2 shows the main thread for the parallel enumeration. It is responsible to initialize the first Sub-ENUM thread and manage the thread list L. A Sub-ENUM thread (SET) is represented by the tuple ( $\mathbf{u}, l, c, t$ ), where $\mathbf{u}$ is the coefficient vector, $l$ the intermediate norm of the root to this subtree, $c$ the search region center and $t$ the lattice dimension minus the starting depth of the parent node in the search tree.

```
Algorithm 2: Main thread for parallel enumeration
    Input: Gram-Schmidt coefficients \(\left(\mu_{i, j}\right)_{1 \leq j \leq i \leq n},\left\|\mathbf{b}_{1}^{*}\right\|^{2} \ldots\left\|\mathbf{b}_{n}^{*}\right\|^{2}\)
    Output: \(\mathbf{u}_{\min }\) such that \(\left\|\sum_{i=1}^{n} u_{i} \mathbf{b}_{i}\right\|=\lambda_{1}(\overline{\mathcal{L}}(\mathbf{B}))\)
    \(A \leftarrow\left\|\mathbf{b}_{1}^{*}\right\|^{2}, \mathbf{u}_{\text {min }} \leftarrow(1,0, \ldots, 0) \quad \triangleright\) Global variables
    \(\mathbf{u} \leftarrow(1,0, \ldots, 0), l \leftarrow 0, c \leftarrow 0, t \leftarrow 1 \quad \triangleright\) Local variables
    \(L \leftarrow\{(\mathbf{u}, l, c, t)\} \quad \triangleright\) Initialize list
    while \(L \neq \emptyset\) or threads are running do
        if \(L \neq \emptyset\) and cores available then
            pick \(\Delta=(\mathbf{u}, l, c, t)\) from L
            start Sub-ENUM thread \(\Delta=(\mathbf{u}, l, c, t)\) on new core
```

Whenever the list contains a SET and free processor units exist, the first SET of the list is executed. The execution of SETs is performed by Algorithm 3. We process the search tree in the same manner as the serial algorithm (Algorithm 1), except the introduction of the loop bound bound and the handling of new SETs (lines $9-11$ ). First, the loop bound controls the termination of the subtree and prohibits that nodes are visited twice. Second, only the SET whose bound is set to the lattice dimension is allowed to create new SETs. Otherwise, if we

```
Algorithm 3: Sub-ENUM thread (SET)
    Input: Gram-Schmidt coefficients \(\left(\mu_{i, j}\right)_{1 \leq j \leq i \leq n},\left\|\mathbf{b}_{1}^{*}\right\|^{2} \ldots\left\|\mathbf{b}_{n}^{*}\right\|^{2},(\overline{\mathbf{u}}, \bar{l}, \bar{c}, \bar{t})\)
    \(\mathbf{u} \leftarrow \overline{\mathbf{u}}, l \leftarrow(0, \ldots, 0), c \leftarrow(0, \ldots, 0)\)
    \(t \leftarrow \bar{t}, l_{t+1} \leftarrow \bar{l}, c_{t} \leftarrow \bar{c}\), bound \(\leftarrow n\)
    while \(t \leq\) bound do
        \(l_{t} \leftarrow l_{t+1}+\left(u_{t}+c_{t}\right)^{2}\left\|\mathbf{b}_{t}^{*}\right\|^{2}\)
        if \(l_{t}<A\) then
                if \(t>1\) then
                \(t \leftarrow t-1 \quad \triangleright\) move one layer down in the tree
                \(c_{t} \leftarrow \sum_{i=t+1}^{n} u_{i} \mu_{i, t}, \quad u_{t} \leftarrow\left\lceil c_{t}\right\rfloor\)
                if bound \(=n\) then
                    \(L \leftarrow L \cup\left(\mathbf{u}, l_{t+2}, c_{t+1}, t+1\right) \quad \triangleright\) insert new \(S E T\) in list \(L\)
                    bound \(\leftarrow t\)
            else
                \(A \leftarrow l_{t}, \quad \mathbf{u}_{\text {min }} \leftarrow \mathbf{u} \quad \triangleright\) set new global minimum
        else
            \(t \leftarrow t+1 \quad \therefore\) move one layer up in the tree
            choose next value for \(u_{t}\) using the zig-zag pattern
```

allow each SET to create new SETs by itself, this would lead to an explosion of the number of threads and each thread has too few computations to perform. We denote the SET with bound set to $n$ by unbounded SET (USET). At any time, there exists only one USET that might be stored in the thread list $L$.

As soon as an USET has the chance to find a new minimum within the current subtree (lines $5-6$ ), its bound is set to the current $t$ value. Thereby, it is transformed to a SET and the new created SET becomes the USET.

### 4.3 Improvements

We presented a first solution for the parallelization of the ENUM algorithm providing a runtime speed-up by a divide and conquer technique. We distribute subtrees to several processor units to search for the minimum. Our improvements deal with the creation of SETs and result in significantly shorter running time. Recall the definitions of Sub-ENUM thread (SET) and unbounded Sub-ENUM thread (USET). By now we call a node, where a new SET can be created, a candidate. Note that a candidate can only be found in an USET.

The following paragraphs show worst cases of the presented parallel ENUM algorithm and present possible solutions to overcome the existing drawbacks.

Threads within threads. Our parallel ENUM algorithm allows to create new SETs only by an USET. The avoidance of producing immense overhead which happens by permitting the creation of new SETs by any SET, backs our decision that it suffices to let only USET create new instances. However, if an USET creates a new SET at a node of depth 1 , then this new SET is executed by a single processor sequentially. Note that this SET solves the SVP problem in dimension $n-1$. It turns out that in the case the depth of a current analyzed node in ENUM is sufficient far away from the depth $t$ of the starting node, the creation
of a new SET is advantageous according to the overall running time and the number of simultaneously occupied processors. Therefore, we introduce a bound $s_{\text {deep }}$ which expresses what we consider to be sufficient far away, i.e. if a SET visits a node with depth $k$ fulfilling the equation $k-t \geq s_{\text {deep }}$ where $t$ stands for the depth of the starting node and it is not an USET, then this SET is permitted to create a new SET once.

Thread Bound. Although we avoid the execution of SETs where the dimension of the subtree is too big, we are still able to optimize the parallel ENUM algorithm by considering execution bounds. We achieve additional performance improvements by the following idea. Instead of generating SETs in each possible candidate, we consider the depth of the node. This enables us to avoid big subtrees for new SETs by introducing an upper bound $s_{\text {up }}$ representing the minimum distance of a node to the root to become a candidate. If ENUM visits a node with depth $t$ fulfilling $n-t>s_{u p}$ and this node is a candidate, we no longer make a subtree ready for a new SET. We rather prefer to behave in that situation like the serial ENUM algorithm. Good choices for the above bounds $s_{\text {deep }}$ and $s_{u p}$ are evaluated in Section 5.

## 5 Experiments

We performed numerous experiments to test our parallel enumeration algorithm. We created 5 different random lattices of each dimension $n \in\{42, \ldots, 56\}$ in the sense of Goldstein and Mayer [GM03]. The bitsize of the entries of the basis matrices were in the order of magnitude of 10 n . We started with bases in Hermite normal form, then LLL-reduced the bases (using LLL parameter $\delta=0.99)$. The experiments were performed on a compute server equipped with four AMD Opteron $(2.3 \mathrm{GHz})$ quad core processors. We compare our results to the highly optimized, serial version of fpLLL in version 3.0.12, the fastest ENUM implementation known, on the same platform. The programs were compiled using gec version 4.3.2. For handling parallel processes, we used the Boost-library in version 1.40. Our implementation uses double precision to store the GramSchmidt coefficients $\mu_{i, j}$ and the $\left\|\mathbf{b}_{1}^{*}\right\|^{2}, \ldots,\left\|\mathbf{b}_{n}^{*}\right\|^{2}$. Due to [PS08], this is suitable up to dimension 90 , which seems to be out of the range of today's enumeration algorithms.

We tested the parallel ENUM algorithm for several $s_{\text {deep }}$ values and concluded that $s_{\text {deep }}=\frac{25}{36}(n-t)$ seems to be a good choice, where $t$ is the depth of the starting node in a SET instance. Further, we use $s_{u p}=\frac{5}{6} n$.

Table 1 and Figure 2 present the experimental results that compare our parallel version to our serial algorithm and to the fpLLL library. We only present the timings, as the output of the algorithms is in all cases the same, namely a shortest non-zero vector of the input lattice. The corresponding speed-ups are shown in Figure 3.

To show the strength of parallelization of the lattice enumeration, we first compare our multi-core versions to our single-core version. The best speed-ups

| $n$ | 42 | 44 | 46 | 48 | 50 | 52 | 54 | 56 |
| :---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| 1 core | 3.81 | 27.7 | 37.6 | 241 | 484 | 3974 | 10900 | 223679 |
| 4 cores | 0.99 | 7.2 | 8.8 | 55 | 107 | 976 | 2727 | 56947 |
| 8 cores | 0.62 | 4.0 | 4.8 | 28 | 56 | 504 | 1390 | 28813 |
| 16 cores | 0.52 | 2.6 | 3.5 | 18 | 36 | 280 | 794 | 16583 |
| fpLLL 1 core | 3.32 | 23.7 | 29.7 | 184 | 367 | 3274 | 9116 | 184730 |

Table 1. Average time in seconds for enumeration of lattices in dimension $n$.


Fig. 2. Average runtimes of enumeration of 5 random lattices in each dimension, comparing our multi-core implementation to fpLLL's and our own single-core version. The right picture shows the data of the left picture in logarithmic scale.
are $4.5(n=50)$ for 4 cores, $8.6(n=50)$ for 8 cores, and $14.2(n=52)$ for 16 cores. This shows that, using $s$ processor cores, we sometimes gain speed-ups of more than $s$, which corresponds to an efficiency of more than 1 . This is a very untypical behavior for parallel algorithms, and is a nice attribute of lattice enumeration. It is caused by the extra communication for the write-back of the current minimum $A$.

The highly optimized enumeration of fpLLL is around $10 \%$ faster than our serial version. Compared to the fpLLL algorithm, we gain a speed-up of up to $6.6(n=48)$ using 8 CPU cores and up to $11.7(n=52)$ using 16 cores. This corresponds to an efficiency of 0.825 ( 8 cores) and 0.73 ( 16 cores), respectively. Therefore, we are beating the best algorithm available with a factor of more than 10 in lattice dimension bigger than 48 .

## 6 Conclusion and Further Work

In this paper we have presented a parallel version of the most common algorithm for solving the shortest vector problem in lattices, the ENUM algorithm. We have shown that a huge speed-up and a high efficiency is reachable using multi-core processors. As parallel versions of LLL are already known, with our parallel ENUM we have given evidence that both parts of the BKZ reduction
algorithm can be parallelized. It remains to combine both, parallel LLL and parallel ENUM, to a parallel version of BKZ. Our experience with BKZ shows that in higher blocksizes of $\approx 50$ ENUM takes more than $99 \%$ of the complete runtime. Therefore, the speed-up of ENUM will directly speed up BKZ reduction, which in turn influences the security of lattice based cryptosystems.

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Fig. 3. Average speed-up of parallel ENUM compared to single-core version (left) and compared to fpLLL single-core version (right).
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[^0]:    ${ }^{3}$ On a 24 core machine we gain speed-ups of 22 ; those experiments will be shown in the final version of this paper.

